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(54) **BIOFUEL COMPOSITION, PROCESS OF PREPARATION AND A METHOD OF FUELING THEREOF**

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**C10L 1/02** (2006.01)

*C10L 1/18* (2006.01)  
*C10L 1/19* (2006.01)  
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CPC ..... **C10L 1/026** (2013.01); *C10L 1/1802* (2013.01); *C10L 1/1824* (2013.01); *C10L 1/19* (2013.01); *Y02E 50/13* (2013.01)  
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USPC ..... 204/157.15; 44/457, 302, 418, 307  
See application file for complete search history.

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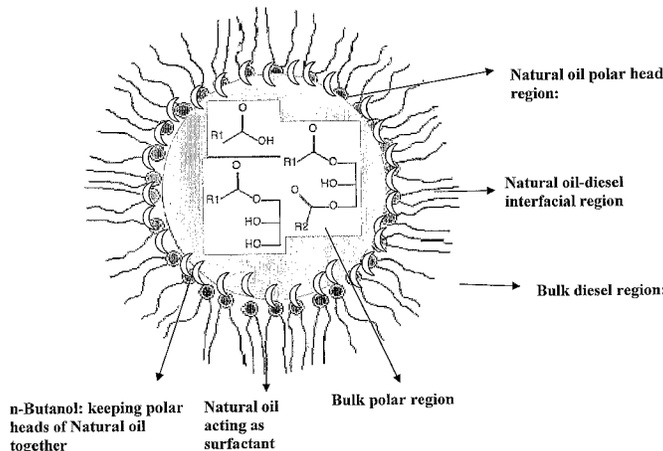
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(57) **ABSTRACT**

The present invention relates to a biofuel composition comprising normal chain alcohol(s), natural oils and diesel optionally along with fuel additive(s); a process of preparation and a method of fueling said biofuel composition.

**25 Claims, 5 Drawing Sheets**



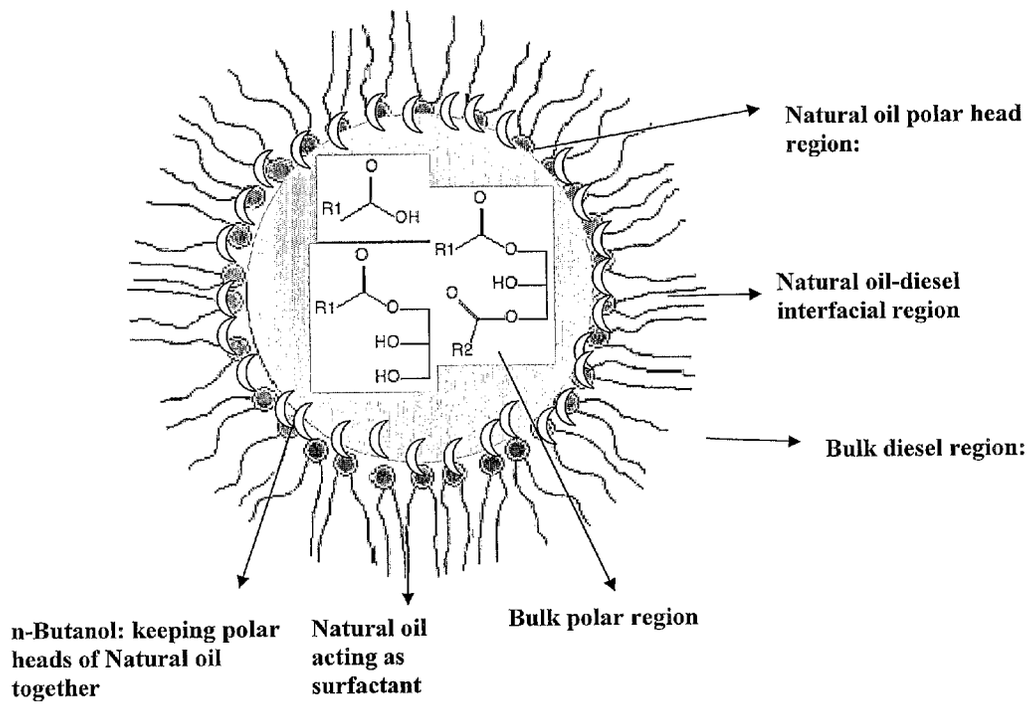


Figure 1

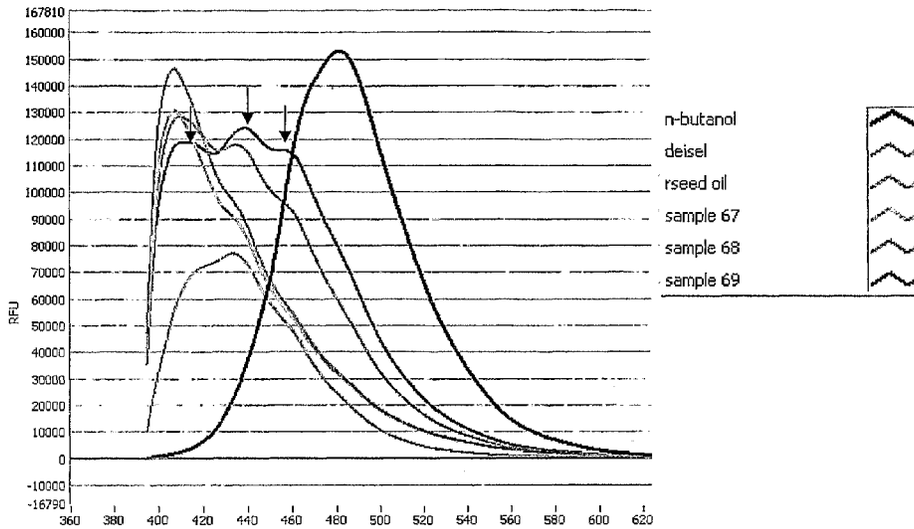


Figure 2

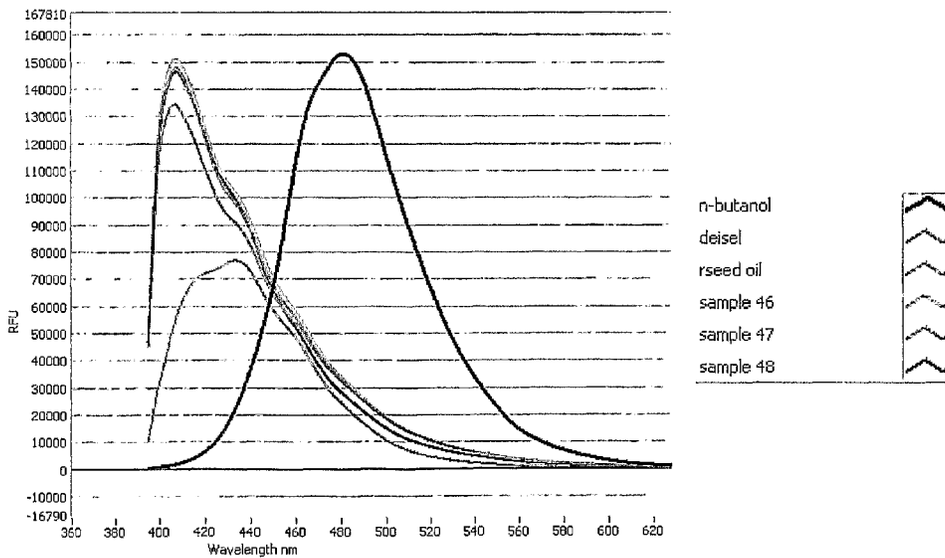


Figure 3

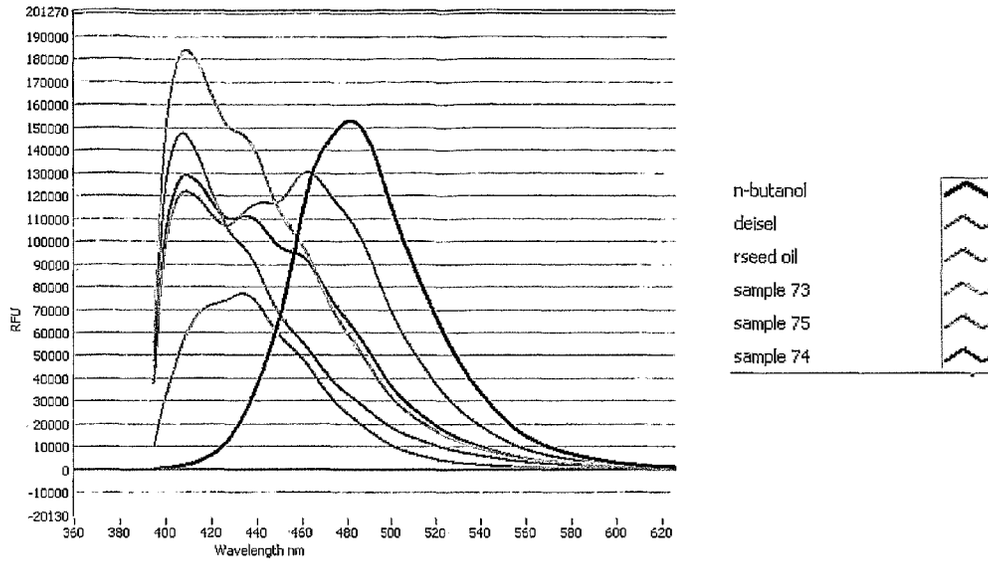


Figure 4

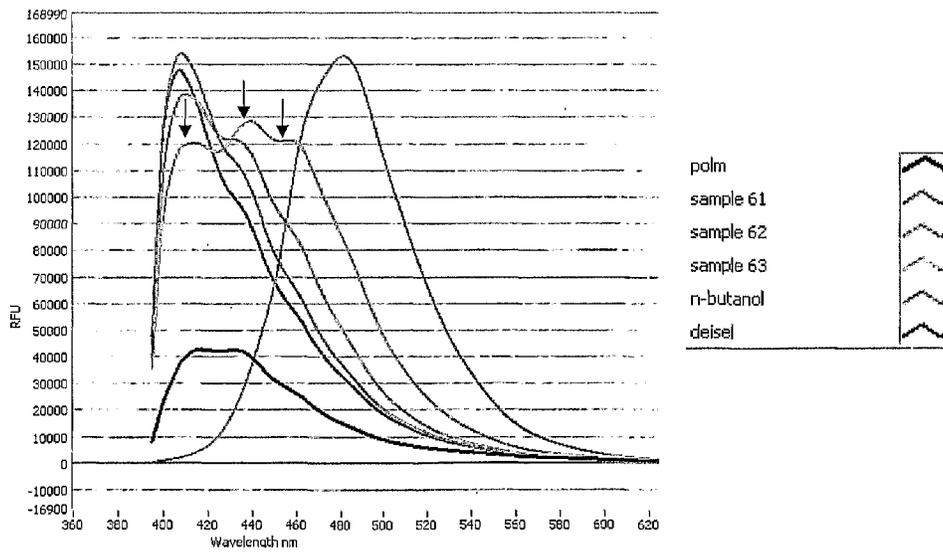


Figure 5

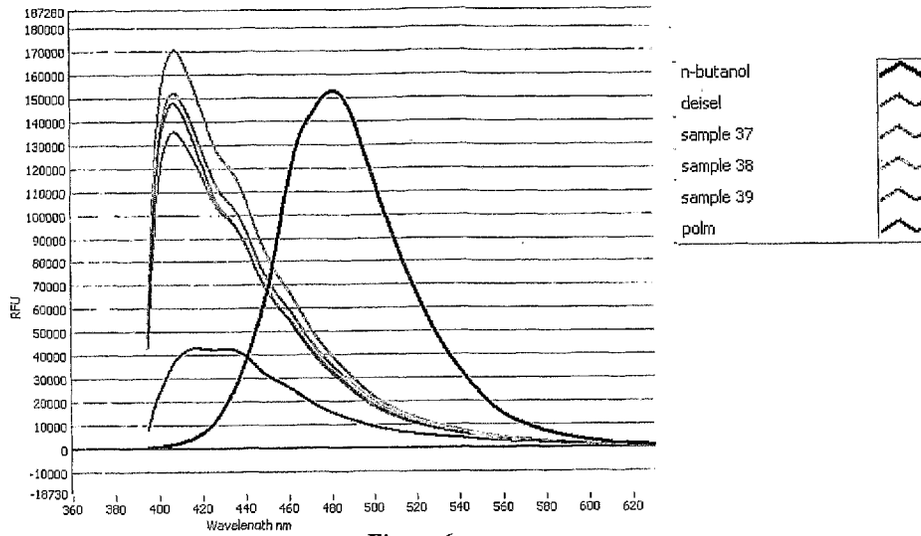


Figure 6

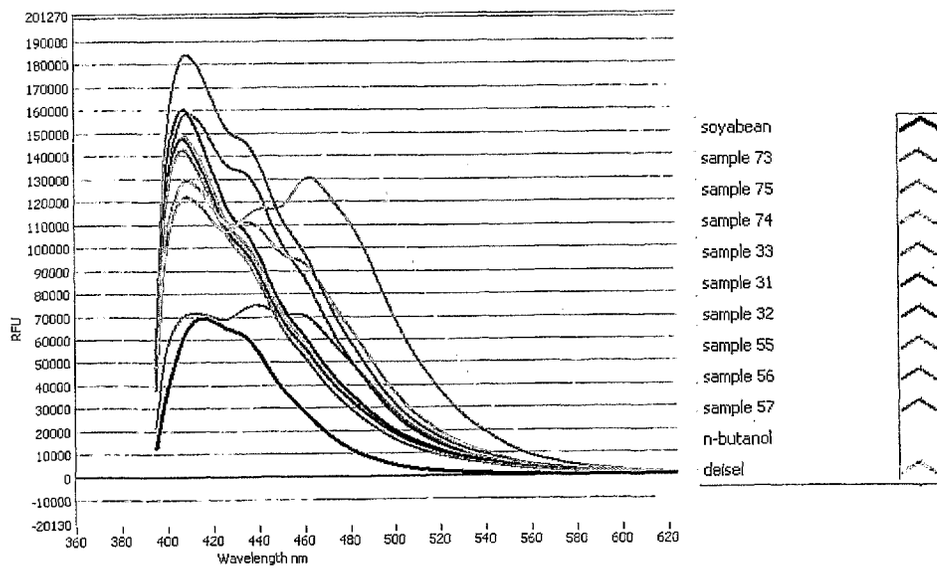


Figure 7

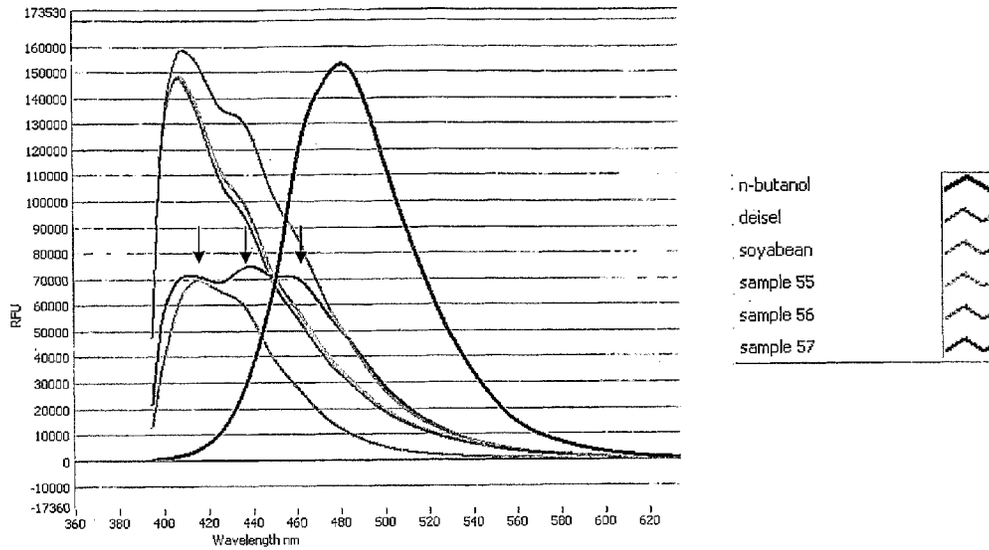


Figure 8

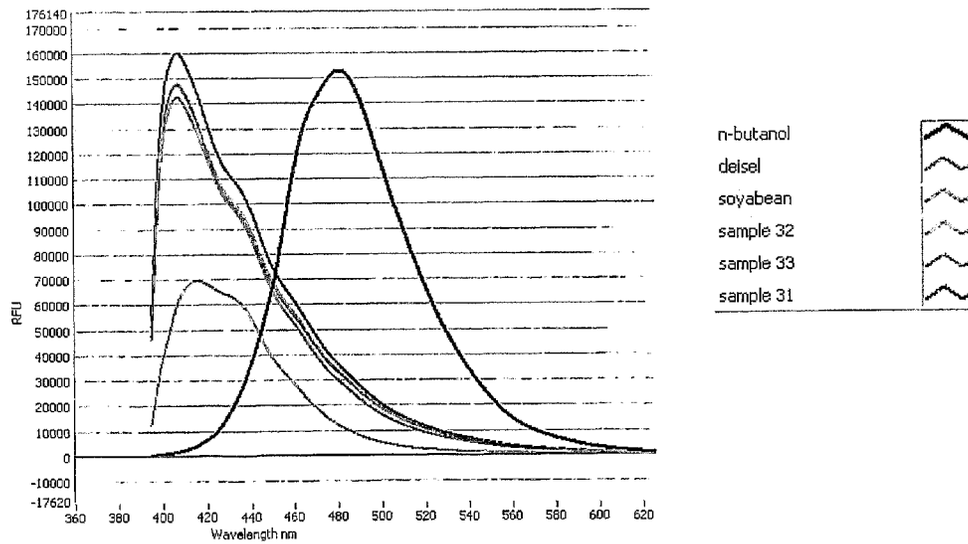


Figure 9

**BIOFUEL COMPOSITION, PROCESS OF PREPARATION AND A METHOD OF FUELING THEREOF**

RELATED APPLICATIONS

This application is a U.S. National Phase of International Application No.: PCT/IN2009/000445, filed Aug. 10, 2009, designating the U.S. and published in English on Apr. 15, 2010 as WO 2010/041266 A1, which claims the benefit of Indian Application No. 02450/CHE/2008, filed Oct. 6, 2008.

FIELD OF THE INVENTION

The present invention relates to biofuel. It involves use of normal chain alcohol(s) to stabilize blends of natural oil(s) and diesel. Therefore, the biofuel composition comprises natural oils, diesel and normal chain alcohols as its components optionally alongwith fuel additives.

BACKGROUND AND PRIOR ART

Pure plant oil (PPO) or tree based oil (TBO) or straight vegetable oil (SVO) are the terms used in the context of neat biodiesel. SVO or TBO or PPO are natural oils which rarely is used as such in commercial diesel engines and their use as blends is also limited.

Natural oils also include any oil obtained from animal source as well (animal fat oil). Plant oils such as SVO or PPO or TBO and animal oils are rich in triglycerides and can be obtained by mechanical, chemical & physical processing from the respective sources.

Natural oils as blends of diesel have some properties that prevent their use in commercial diesel engines. The major ones are listed below.

1. High boiling point: Natural oils boil at higher temperature than diesel and have problems meeting the norms required for minimum distillation characteristics. This causes the residue formation in engine and in long run there will be accumulation of greasy material in the engine. This could lead to clogging, increased smoke at higher loads, increase in emission of nitrogen oxides (NOx), loss of power and mileage, sudden stoppage of engine.
2. High viscosity: Natural oils are about eight times viscous than diesel and as such cannot be used in current diesel engines without modification. Natural oil blends of diesel (below 20%) often meet the physico chemical properties of diesel and were used without any short-term issues. But in long term, they cause non-uniform spraying at ignition chamber, engine deposits, inefficient burning of fuel and increase in NOx.
3. Immiscibility: Natural oils are miscible with diesel and stay miscible for 15-30 days depending on the composition of oil, but longer periods of storage cause separation of natural oils and diesel layers and in about two months storage, almost complete separation occurs. The separated layer looks like a brown or yellow precipitate and this causes engine deposits, sudden stoppage/malfunction of engine, increase in emission of particulate matter, etc.

The main issues with respect to diesel engine by use of blends of natural oils with diesel are

Issues	Reason	Possible solution
a) Clogging	Deposits and greasy material forming in engine	Grease or engine deposit formation need to be prevented (transesterification of natural oil is one method)
b) Increase in smoke	Improper mixing of natural oils and diesel, gradual separation of layers	Miscibility with diesel need to be improved (by adding some additives)
c) Increase in NOx	Any strain on fuel delivery/burning causes increase in NOx. Strain in fuel delivery is mainly caused by non-uniformity of fuel.	Fuel injected need to be uniform in composition and quality
d) Loss of power and mileage	Even though natural oils have rich oxygen content and higher calorific value than diesel, loss of power and mileage occur due to non-uniformity of fuel.	Separation of natural oils and diesel layers is the main cause and natural oils needs to be made completely miscible with diesel
e) Sudden stoppage of engine	Increase in moisture content in engine due to the waxy solid deposition attracts dust and more moisture that are retained inside the engine. Eventually, these enter the ignition chamber causing abrupt change in fuel behavior causing sudden stoppage of engine.	The waxy or grease solid formation need to be avoided and any free fatty acids or partially hydrolysed triglycerides present in natural oils need to be solubilized in diesel.
f) Engine deposits	Separation of natural oils and diesel and their in turn assistance in accumulation of dust and moisture is responsible for engine deposits.	Natural oils (SVO) are highly viscous liquids with minute percentage of free fatty acids or partially hydrolysed triglycerides (diglycerides, monoglycerides) produced during mechanical or solvent based processing of seeds/bark/leaves. These free fatty acids precipitate from blended diesel within one day and increase the engine deposit formation and somehow these fatty acids need to be solubilized in diesel.

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Issues	Reason	Possible solution
g) Increase in sound from engine	Engine deposits and non uniformity of fuel flow results in stress on compression ignition system, which results in increased noise	Elimination of any insoluble particle in blended diesel and ensuring uniform flow of fuel.

It is clearly understood from above table that most of the problems caused by natural oil blends with diesel is immiscibility. When the natural oil has the property of not being soluble in diesel, the above problems like engine deposits, increase in NOx emission, increased smoke, decrease in power and mileage, etc occur.

To address the problems associated with use of natural oil in diesel, biodiesel manufacturers use transesterified oil obtained through transesterification reaction. This way the triglycerides are converted to free fatty acids and the fatty acids are in turn esterified with methanol or ethanol (use of higher alcohols is also reported). The hydrolysis process involves use of sodium hydroxide and methanol and is a heat intensive process (lot of heat from environment to be supplied). There is increasing criticism that whatever benefit derived by shifting to renewable plant based oils (reduction in carbon dioxide, lower pollution, etc), is being lost by use of toxic industrial chemicals (like methanol), problems with waste disposal, increased heat input etc.

Hence, there is a need for development of methods to avoid transesterification of natural oils. A process to convert the natural oils to a diesel blend product is credible, but identifying compounds or components which can stabilize natural oil blends will be of great interest and could provide a cheaper, safer and environmentally friendly alternative to transesterification, help in reduction of pollution, carbon dioxide emissions, decrease in byproduct toxicity, no heat input from environment, etc.

PCT/US99/00598 discloses an additive composition consisting of water soluble alcohols. All the composition disclosed in PCT/US99/00598 shows the presence of water and ethanol.

Yoshimoto et al (Published by society of automotive engineers, 400 commonwealth drive, Warrendale, Pa.-15096, USA, October 2002) discloses performance of a diesel engine by the blends of rapeseed oil and oxygenated organic compounds such as ethanol, 1-propanol, 1-butanol, 1-pentanol, 2-methoxyethanol, 2-ethoxyethanol, 2-butoxyethanol and dibutyl ether.

Khaled et al (1<sup>st</sup> International energy conversion engineering conference, Virginia, USA, August 2003) discloses fuel blends in compression ignition engines which fuel blend comprises iso-butanol, diesel and olive oil.

The present invention addresses the problems associated with prior art mentioned above.

#### OBJECTS OF THE INVENTION

The main object of the present invention is to obtain a biofuel composition comprising normal chain alcohol(s), natural oils and diesel.

Another object of the present invention is to develop a process for obtaining the biofuel composition. biofuel composition.

#### STATEMENT OF THE INVENTION

Accordingly, the present invention relates to a biofuel composition comprising normal chain alcohol(s) having greater

than 3 carbon atoms but not more than 21 carbon atoms at a concentration ranging from about 1% to about 24% v/v, natural oils at a concentration ranging from about 1% to about 24% v/v and diesel at a concentration ranging from about 52% to about 98% v/v optionally alongwith fuel additive(s); a biofuel composition comprising n-butanol at a concentration ranging from about 1% to about 24% v/v, natural oils at a concentration ranging from about 1% to about 24% v/v and diesel at a concentration ranging from about 52% to about 98% v/v optionally alongwith fuel additive(s); a process for preparation of biofuel composition comprising normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration ranging from about 1% to about 24% v/v, natural oils at a concentration ranging from about 1% to about 24% v/v and diesel at a concentration ranging from about 52% to about 98% v/v optionally alongwith fuel additive(s), said process comprises mixing said normal chain alcohol(s) with a blend comprising the natural oils and the diesel optionally alongwith fuel additive(s); and a method of fueling biofuel composition comprising normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration ranging from about 1% to about 24% v/v, natural oils at a concentration ranging from about 1% to about 24% v/v and diesel at a concentration ranging from about 52% to about 98% v/v optionally alongwith fuel additive(s), said method comprising fueling the biofuel composition for fueling applications.

#### BRIEF DESCRIPTION OF ACCOMPANYING FIGURES

FIG. 1 Reverse micelle structure in natural oil blended diesel in presence of normal chain alcohols. Reverse micelle in diesel shows bulk diesel region, natural oil-diesel interfacial region, natural oil polar head region and bulk polar region. Natural oil polar head region: This region has n-butanol in combination with triglyceride polar groups. n-butanol helps the polar heads to form stable reverse micelles by increasing polarity of system; Natural oil-diesel interfacial region: This is formed by addition of n-butanol (n-butanol, natural oil and diesel); Bulk diesel region: Almost diesel characteristics, very non polar; Bulk polar region: Polar components like free fatty acids, monoglycerides and diglycerides produced from degradation of natural oil could be present. This region could also hold traces of water absorbed by the polar components over time, preventing engine deposit formation.

FIG. 2 shows the existence of reverse micelle in fuel blends containing rapeseed oil, n-butanol and diesel. The samples are n-butanol, diesel, rapeseed oil (SVO), Sample 67 (5% rapeseed oil, 5% n-butanol, 90% diesel), Sample 68 (10% rapeseed oil, 10% n-butanol, 80% diesel), Sample 69 (20% rapeseed oil, 20% n-butanol, 60% diesel). Samples 67, 68 and 69 show 3 peaks corresponding to 3 regions in reverse micelles (bulk diesel region, SVO-diesel interfacial region and SVO polar head region).

FIG. 3 shows the absence of reverse micelle in fuel blends containing natural oil and diesel. The samples are n-butanol, diesel, rapeseed oil (rseed oil, SVO), Sample 46 (10% rapeseed oil, 90% diesel), Sample 47 (20% rapeseed oil, 80% diesel), Sample 48 (40% rapeseed oil, 60% diesel). It can be clearly observed that, without n-butanol, the samples have behavior similar to diesel and after standing for few days, due to absence of stable interactions of polar heads of rapeseed oil with diesel, the rapeseed oil separates from diesel.

FIG. 4 shows that n-butanol has the property of formation of reverse micelle in diesel and existence of three different

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regions. This is n-butanol control for all the oils in the study. The samples are n-butanol, diesel, rapeseed oil (rseed oil, SVO), Sample 73 (10% n-butanol, 90% diesel), Sample 74 (20% n-butanol, 80% diesel), Sample 75 (40% n-butanol, 60% diesel). n-butanol blends in diesel also show three regions similar to natural oil and n-butanol blends with diesel. This indicates that the reverse micellar structure is originating from presence of n-butanol and it is aiding natural oil to form similar structures in diesel.

FIG. 5 shows the existence of reverse micelle in fuel blends containing palmoline oil, n-butanol and diesel. The samples are n-butanol, diesel, palmoline oil (polm, SVO), Sample 61 (5% palmoline oil, 5% n-butanol, 90% diesel), Sample 62 (10% palmoline oil, 10% n-butanol, 80% diesel), Sample 63 (20% palmoline oil, 20% n-butanol, 60% diesel). N-butanol blends in diesel show three regions similar to natural oil and n-butanol blends with diesel. This indicates that the reverse micellar structure is originating from presence of n-butanol and it is aiding natural oil to form similar structures in diesel. Samples 61, 62 and 63 shows 3 peaks corresponding to 3 regions in reverse micelles (bulk diesel region, SVO-diesel interfacial region and SVO polar head region).

FIG. 6 shows the absence of reverse micelle in fuel blends containing SVO and diesel. The samples are n-butanol, diesel, palmoline oil (polm, SVO), Sample 37 (10% palmoline oil, 90% diesel), Sample 38 (20% palmoline oil, 80% diesel), Sample 39 (40% palmoline oil, 60% diesel). It can be clearly observed that, without n-butanol, the samples have behavior similar to diesel and after standing for few days, due to absence of stable interactions of polar heads of SVO with diesel, the SVO separates from diesel.

FIG. 7 shows the study on soyabean oil (SVO) using PRODAN. The samples are n-butanol, diesel, soyabean oil (SVO), Sample 55 (5% soyabean oil, 5% n-butanol, 90% diesel), Sample 56 (10% soyabean oil, 10% n-butanol, 80% diesel), Sample 57 (20% soyabean oil, 20% n-butanol, 60% diesel), Sample 74 (20% n-butanol, 80% diesel), Sample 73 (10% n-butanol, 90% diesel), Sample 75 (40% n-butanol, 60% diesel), Sample 31 (10% soyabean oil, 90% diesel), Sample 32 (20% soyabean oil, 90% diesel), Sample 33 (40% soyabean oil, 60% diesel)

FIG. 8 shows the existence of reverse micelle in fuel blends containing soyabean oil, n-butanol and diesel. The samples are n-butanol, diesel, soyabean oil (SVO), Sample 55 (5% soyabean oil, 5% n-butanol, 90% diesel), Sample 56 (10% soyabean oil, 10% n-butanol, 80% diesel), Sample 57 (20% soyabean oil, 20% n-butanol, 60% diesel). Samples 55, 56 and 57 shows 3 peaks corresponding to 3 regions in reverse micelles (bulk diesel region, SVO-diesel interfacial region, and SVO polar head region).

FIG. 9 shows the absence of reverse micelle in fuel blends containing soyabean oil and diesel. The samples are n-butanol, diesel, soyabean oil (SVO), Sample 31 (10% soyabean oil, 90% diesel), Sample 32 (20% soyabean oil, 80% diesel), Sample 33 (40% soyabean oil, 60% diesel). It can be clearly observed that, without n-butanol, the samples have behavior similar to diesel and after standing for few days, due to absence of stable interactions of polar heads of soyabean oil with diesel, the soyabean oil separates from diesel.

#### DETAILED DESCRIPTION OF THE INVENTION

The present invention relates to a biofuel composition comprising normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration ranging from about 1% to about 24% v/v, natural oils at a concentration ranging from about 1% to about 24% v/v

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and diesel at a concentration ranging from about 52% to about 98% v/v optionally alongwith fuel additive(s).

In another embodiment of the present invention, the normal chain alcohols are selected from a group comprising n-butanol, n-pentanol, n-hexanol, n-heptanol, n-octanol (capryl alcohol), n-nonanol (pelargonic alcohol), n-decanol (capric alcohol), n-dodecanol (lauryl alcohol), n-pentadecanol, n-hexadecanol (cetyl alcohol), n-tetradecanol (myristyl alcohol), cis-9-hexadecen-1-ol (palmitoleyl alcohol), n-octadecanol (stearyl alcohol), 9E-octadecen-1-ol (elaidyl alcohol), cis-9-octadecen-1-ol (oleyl alcohol), 9Z, 12Z-octadecadien-1-ol (linoleyl alcohol), 9E, 12E-octadecadien-1-ol (elaidolinoleyl alcohol), 9Z, 12Z, 15Z-octadecatrien-1-ol (linolenyl alcohol), 9E, 12E, 15-E-octadecatrien-1-ol (elaidolinolenyl alcohol), 12-hydroxy-9-octadecen-1-ol (ricinoleyl alcohol) and 1-eicosanol (arachidyl alcohol) or any combinations thereof.

In yet another embodiment of the present invention, the normal chain alcohol is preferably n-butanol.

In still another embodiment of the present invention, the natural oils are selected from a group comprising straight vegetable oil (SVO), plant parts oils (PPO), tree based oil (TBO), animal fat oil or any combinations thereof.

In still another embodiment of the present invention, the ratio of normal chain alcohol(s) and natural oils is preferably 1:1.

In still another embodiment of the present invention, the fuel additive(s) is selected from a group comprising anti-polymerizing agents, anti-knocking agents anti-freezing agents, anti-coagulation agents, anti-precipitation agents, cetane index boosters and cetane number boosters or any combinations thereof.

In still another embodiment of the present invention, the fuel additive(s) is present at a concentration ranging from about 0.01 to about 5% v/v, preferably about 0.01% to about 1% v/v.

In still another embodiment of the present invention, the biofuel composition comprises normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 5% v/v, natural oils at a concentration of about 5% v/v and diesel at a concentration of about 90% v/v optionally alongwith fuel additive(s).

In still another embodiment of the present invention, the biofuel composition comprises normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 7.5% v/v, natural oils at a concentration of about 7.5% v/v and diesel at a concentration of about 85% v/v optionally alongwith fuel additive(s).

In still another embodiment of the present invention, the biofuel composition comprises normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 10% v/v, natural oils at a concentration of about 10% v/v and diesel at a concentration of about 80% v/v optionally alongwith fuel additive(s).

In still another embodiment of the present invention, the biofuel composition comprises normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 20% v/v, natural oils at a concentration of about 20% v/v and diesel at a concentration of about 60% v/v optionally alongwith fuel additive(s).

In still another embodiment of the present invention, the biofuel composition comprises normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 10% v/v, natural oils at a concentration of about 5% v/v and diesel at a concentration of about 85% v/v optionally alongwith fuel additive(s).

The present invention relates to a biofuel composition comprising n-butanol at a concentration ranging from about 1% to about 24% v/v, natural oils at a concentration ranging from about 1% to about 24% v/v and diesel at a concentration ranging from about 52% to about 98% v/v optionally along-

with fuel additive(s).  
 The present invention relates to a process for preparation of biofuel composition comprising normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration ranging from about 1% to about 24% v/v, natural oils at a concentration ranging from about 1% to about 24% v/v and diesel at a concentration ranging from about 52% to about 98% v/v optionally alongwith fuel additive(s), said process comprises mixing said normal chain alcohol(s) with a blend comprising the natural oils and the diesel optionally alongwith fuel additive(s).

In still another embodiment of the present invention, the normal chain alcohols are selected from a group comprising n-butanol, n-pentanol, n-hexanol, n-heptanol, n-octanol (capryl alcohol), n-nonanol (pelargonic alcohol), n-decanol (capric alcohol), n-dodecanol (lauryl alcohol), n-pentadecanol, n-hexadecanol (cetyl alcohol), n-tetradecanol (myristyl alcohol), cis-9-hexadecen-1-ol (palmitoleyl alcohol), n-octadecanol (stearyl alcohol), 9E-octadecen-1-ol (elaidyl alcohol), cis-9-octadecen-1-ol (oleyl alcohol), 9Z, 12Z-octadecadien-1-ol (linoleyl alcohol), 9E, 12E-octadecadien-1-ol (elaidolinoleyl alcohol), 9Z, 12Z, 15Z-octadecatrien-1-ol (linolenyl alcohol), 9E, 12E, 15-E-octadecatrien-1-ol (elaidolinolenyl alcohol), 12-hydroxy-9-octadecen-1-ol (ricinoleyl alcohol) and 1-eicosanol (arachidyl alcohol) or any combinations thereof.

In still another embodiment of the present invention, the natural oils are selected from a group comprising straight vegetable oil (SVO), plant parts oils (PPO), tree based oil (TBO), animal fat oil or any combinations thereof.

In still another embodiment of the present invention, the natural oils are obtained by mechanical or chemical treatment of the source from which it is derived.

In still another embodiment of the present invention, the ratio of normal chain alcohol(s) and natural oils is preferably 1:1.

In still another embodiment of the present invention, the fuel additive(s) is selected from a group comprising anti-polymerizing agents, anti-knocking agents anti-freezing agents, anti-coagulation agents, anti-precipitation agents, cetane index boosters and cetane number boosters or any combinations thereof.

In still another embodiment of the present invention, the fuel additive(s) is present at a concentration ranging from about 0.01 to about 5% v/v, preferably about 0.01% to about 1% v/v.

In still another embodiment of the present invention, the process provides for biofuel composition comprising normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 5% v/v, natural oils at a concentration of about 5% v/v and diesel at a concentration of about 90% v/v optionally alongwith fuel additive(s).

In still another embodiment of the present invention, the process provides for biofuel composition comprising normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 7.5% v/v, natural oils at a concentration of about 7.5% v/v and diesel at a concentration of about 85% v/v optionally alongwith fuel additive(s).

In still another embodiment of the present invention, the process provides for biofuel composition comprising normal

chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 10% v/v, natural oils at a concentration of about 10% v/v and diesel at a concentration of about 80% v/v optionally alongwith fuel additive(s).

In still another embodiment of the present invention, the process provides for biofuel composition comprising normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 20% v/v, natural oils at a concentration of about 20% v/v and diesel at a concentration of about 60% v/v optionally alongwith fuel additive(s).

In still another embodiment of the present invention, the process provides for biofuel composition comprising normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 10% v/v, natural oils at a concentration of about 5% v/v and diesel at a concentration of about 85% v/v optionally alongwith fuel additive(s).

In still another embodiment of the present invention, the process provides for biofuel composition comprising butanol at a concentration ranging from about 1% to about 24% v/v, natural oils at a concentration ranging from about 1% to about 24% v/v and diesel at a concentration ranging from about 52% to about 98% v/v optionally alongwith fuel additive(s).

The present invention relates to a method of fueling biofuel composition comprising normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration ranging from about 1% to about 24% v/v, natural oils at a concentration ranging from about 1% to about 24% v/v and diesel at a concentration ranging from about 52% to about 98% v/v optionally alongwith fuel additive(s), said method comprising fueling the biofuel composition for fueling applications.

In still another embodiment of the present invention, the fueling applications are selected from a group comprising space heating, lighting, cooking, running of engines in automobiles, running of motor pumps and generators.

The present invention relates to use of normal-chain alcohols (the alcohol group —OH attached to the terminal carbon) having greater than 3 carbon atoms upto 21 carbon atoms to stabilize natural oil blends in diesel to obtain a biofuel composition. The normal chain alcohols include n-butanol, n-pentanol, n-hexanol, n-heptanol, n-octanol (capryl alcohol), n-nonanol (pelargonic alcohol), n-decanol (capric alcohol), n-dodecanol (lauryl alcohol), n-pentadecanol, n-hexadecanol (cetyl alcohol), n-tetradecanol (myristyl alcohol), cis-9-hexadecen-1-ol (palmitoleyl alcohol), n-octadecanol (stearyl alcohol), 9E-octadecen-1-ol (elaidyl alcohol), cis-9-octadecen-1-ol (oleyl alcohol), 9Z, 12Z-octadecadien-1-ol (linoleyl alcohol), 9E, 12E-octadecadien-1-ol (elaidolinoleyl alcohol), 9Z, 12Z, 15Z-octadecatrien-1-ol (linolenyl alcohol), 9E, 12E, 15-E-octadecatrien-1-ol (elaidolinolenyl alcohol), 12-hydroxy-9-octadecen-1-ol (ricinoleyl alcohol) and 1-eicosanol (arachidyl alcohol) preferably n-butanol or combinations thereof. Normal chain alcohols may be saturated or unsaturated.

The biofuel composition comprises normal chain alcohol(s) having greater than 3 carbon atoms upto 21 carbon atoms ranging from about 1% to about 24%, natural oils ranging from about 1% to about 24% and diesel at a concentration ranging from about 52% to about 98% optionally along with fuel additives. The natural oil used in the present invention may be either crude or refined. The fuel additives are those which are normally added to any diesel fuel composition.

n-butanol can be produced by microbial fermentation, chemically synthesized or obtained from a plant source by

bacterial action. This includes obtaining butanol from cellulose containing plants, lignin-containing plants, from sewage and animal waste, from sugars obtained from plant source and then by fermentation involving algae. Higher alcohols can also be obtained in similar manner and used in the composition. n-butanol is a smallest carbon containing alcohol which can be used for this purpose.

n-dodecanol (lauryl alcohol), n-pentadecanol, n-hexadecanol (cetyl alcohol), n-tetradecanol (myristyl alcohol), cis-9-hexadecen-1-ol (palmitoleyl alcohol), n-octadecanol (stearyl alcohol), 9E-octadecen-1-ol (elaidyl alcohol), cis-9-octadecen-1-ol (oleyl alcohol), 9Z, 12Z-octadecadien-1-ol (linoleyl alcohol), 9E, 12E-octadecadien-1-ol (elaidolinoleyl alcohol), 9Z, 12Z, 15Z-octadecatrien-1-ol (linolenyl alcohol), 9E, 12E, 15-E-octadecatrien-1-ol (elaidolinolenyl alcohol), 12-hydroxy-9-octadecen-1-ol (ricinoleyl alcohol), 1-eicosanol (arachidyl alcohol) can be produced from chemical derivatization of triglycerides. n-octanol (capryl alcohol), n-nonanol (pelargonic alcohol), n-decanol (capric alcohol) can also be produced from plant sources.

Natural oils include straight vegetable oil (SVO), Pure plant oil (PPO), tree based oil (TBO), animal fat oil etc. These are rich in triglycerides and can be obtained by mechanical, chemical & physical processing.

Straight vegetable oil (SVO) is defined as oil obtained from any plant part like, leaves, seeds, bark, flowers, etc. PPO is considered to be equivalent of SVO and includes any oil obtained from a plant source or plant part, including oils obtained from algae.

SVO or PPO can be obtained by mechanical pressing of plant part (eg: crushing and pressing of neem seeds), mechanical processing of plant part (eg: processing of Jatropa seeds to obtain oil), solvent extraction of plant oil (eg: neem seed oil (SVO) obtained by hexane extraction), crushing of leaves (eg: eucalyptus oil), extraction or crushing of wood (eg: cedarwood oil), processing of lipid containing microorganisms (eg: oil from algae) etc.

SVO or PPO obtained from any plant source (edible or non edible) can be used in the biofuel composition of the present invention. SVO or PPO can be obtained from any edible or non-edible part of a plant source including seeds of Jatropa, Pongamia, neem, sunflower, rapeseed, eucalyptus leaves, cedarwood, rose wood, etc.

Minimal chemical processing is defined as the processing involving acids, bases, and alcohols and physical processing involves crushing, pressing, grinding, squeezing etc where mechanical energy is used to obtain the oil.

In general, SVO and PPO are also tree-based oil (TBO) and it should be noted that TBO could be used as part of this invention. In this document, it should be noted that TBO, SVO and PPO are used inter changeably but all of them mean that it is an oil produced from a natural source and is mechanically or chemically treated. Chemical treatment means using an industrial chemical to extract the oil in a chemically modified form. Thus, waste or used natural oils are also used in the composition of present invention.

Chemical treatments include extraction with an organic solvent, water, or distillation or any aqueous based azeotropic distillation.

In this document, the use of words "precipitate" and "no precipitate" with respect to some examples and their definitions are given below.

No precipitate: No visible turbidity, precipitate, separation of layers, settling of greasy material, settling of viscous liquid, droplet formation on the walls or any other evidence of non-uniformity in the liquid.

Precipitate: Observation of turbidity, precipitate, separation of layers, settling of greasy material, two different layers, settling of viscous liquid, droplet formation on the walls or observation of non-uniformity in solution.

In the composition of present invention, any alcohol having greater than 3 carbon upto 21 carbon stabilize blends of natural oil and diesel.

Mechanism of stabilization of natural oil in diesel by normal chain alcohols is by one of the following methods:

a) Increase solubility of natural oil in diesel

Alcohols containing greater than 3 carbons are usually chemical compounds soluble in diesel and natural oil. Being soluble in both (diesel and natural oil), allows them to increase solubility of natural oil in diesel. Due to their higher boiling points (>100° C.), their composition in diesel stays same under normal storage conditions. This allows the composition to remain more or less same for longer storage periods, which results in stable blends with longer shelf life.

b) Formation of non ionic-reverse micelles kind of structure in diesel

When surfactants assemble in oil, the aggregate is referred to as a reverse micelle. In a reverse micelle, the heads are in the core and the tails maintain favorable contact with diesel. Surfactants are also often classified into four primary groups; anionic, cationic, non-ionic and zwitterionic (dual charge).

The above discussed normal chain alcohols tend to aid natural oil to act as surfactants. When natural oil act as surfactants, the non polar tails maintain a favorable interaction with diesel and the polar functional groups form a core which helps in solubilizing any by products formed during the storage of natural oil. Natural oils are known to give out free fatty acids, monoglycerides and diglycerides on storage (in small percentage), which are highly insoluble in diesel and form a greasy material. This is one of the reasons why natural oils are not used as blends of diesel. When fuel additives like n-butanol are used, they tend to increase the solubility of natural oil in diesel and help them act as surfactants which can form reverse micelles. Reverse micelle formation is characterized by observing different fluorescence signals for charge transfer fluorescent probes. The widely used charge transfer fluorescent probe to characterize reverse micelles is PRODAN. Formation of reverse micelle in diesel results in 4 regions namely (FIG. 1):

Bulk diesel region: This region is formed when the diesel is excess of natural oil. This region is characterized by emission in charge transfer fluorescent probes with lower fluorescent emissions.

Natural oil-diesel interfacial region: In this region, Natural oil and diesel are at interface and have fluorescent spectra of charge transfer probes red shifted compared to bulk diesel region.

Natural oil polar head region: Natural oil in diesel form a core of polar functional groups, which can be characterized by further red shift in charge transfer probe's emission spectrum. This is a very polar region and usually the red shift is significant. Importantly, this is the region which will stabilize the formed polar by products of Natural oil and provide long-term stability for Natural oil in diesel.

Bulk polar region: Difficult to observe in current system, as components vary overtime. Mostly polar components like free fatty acids, monoglycerides and diglycerides produced from degradation of Natural oil may be present. This region can hold traces of water absorbed by these polar components. In reverse micelles, this region can be characterized as bulk water region, but the polar impurities possible in natural

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oil-diesel blends are not that polar and hence, the fluorescent peak corresponding to this region could overlap with the natural oil polar head region.

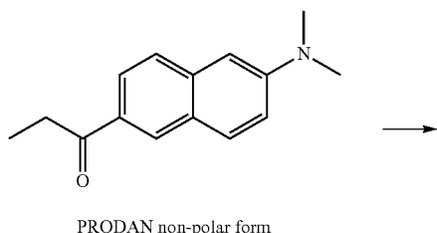
It is observed that a 20% mixture of n-butanol and 20% of natural oils can stay solubilized in diesel for days together indicating that when natural oil is blended with diesel, the by products formed during storage like free fatty acids, monoglycerides, traces of glycerol and diglycerides will be soluble in diesel.

Use of Fluorescent Probes to Detect Reverse Micelle Regions:

A solution is recognized by uniform distribution of solutes in a solvent. Reverse micelles have a unique structure, in which using a charge transfer probe, at least three different regions can be identified.

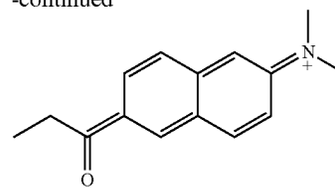
In natural oil/normal chain alcohols/diesel reverse micelle system, the possible regions are bulk diesel region, natural oil-diesel interfacial region, natural oil polar head region and bulk polar region. The surfactant behavior may not be solely from normal chain alcohols or natural oil, but because of combination of both these components. Importantly, the key three regions were identified when a charge transfer fluorescent probe is used.

Charge transfer fluorescent probes like PRODAN exist in two different forms depending on whether it is in polar or non-polar medium. Depending on the polarity of the medium and the ratio of charge transfer, species exist in solution.



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PRODAN polar form

The existence of reverse micelle in fuel blends containing natural oil, normal chain alcohols and diesel of the present invention is shown in FIGS. 2, 5 and 7 with specific examples. From the FIGS. 3, 6 and 9 it is clear that, without normal chain alcohols, the samples have behavior similar to diesel and after standing for few days, due to absence of stable interactions of polar heads of natural oils with diesel, the natural oil separates from diesel. Normal chain alcohol blends in diesel (FIG. 4) also show three regions similar to natural oil and n-butanol blends with diesel. This indicates that the reverse micellar structure is originating from presence of n-butanol and it is aiding natural oil to form similar structures in diesel.

Establishing n-Butanol for Stabilizing Neem Seed Oil Blends with Diesel

Sample details are

	Percentage of diesel	Percentage of Neem seed oil	Percentage of n-butanol	Percentage of t-butanol
Sample 1	100	—	—	—
Sample 2	—	100	—	—
Sample 3	—	—	100	—
Sample 4	—	—	—	100
Sample 6	95	5	—	—
Sample 7	90	10	—	—
Sample 8	90	5	5	—
Sample 9	85	10	5	—
Sample 10	90	5	—	5
Sample 11	85	10	—	5

Results of the experiment are

	0 Days	1 Days	7 Days	14 Days	30 Days	90 Days
Sample 1	No precipitate					
Sample 2	No precipitate					
Sample 3	No precipitate					
Sample 4	No precipitate					
Sample 6	No precipitate	Precipitate	Precipitate	Precipitate	Precipitate	Precipitate
Sample 7	Precipitate	Precipitate	Precipitate	Precipitate	Precipitate	Precipitate
Sample 8	No precipitate					
Sample 9	No precipitate					
Sample 10 (t-butanol)	No precipitate	No precipitate	No precipitate	Precipitate	Precipitate	Precipitate
Sample 11 (t-butanol)	No precipitate	No precipitate	Precipitate	Precipitate	Precipitate	Precipitate

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The above experiment clearly concludes that n-butanol can be used for stabilizing neem seed oil (SVO) blends with diesel. Though t-butanol has the same number of carbons as n-butanol, it fails to stabilize neem seed oil (SVO) blend with diesel for more than 7 days, clearly demonstrating that linear carbon chain of n-butanol played a role. This also establishes that any alcohol, which is less polar than n-butanol (like alcohols containing higher carbons than n-butanol) can be used for this purpose.

The table shows that n-butanol has highest stability as compared to iso-butanol and tertiary butanol.

Role of Linear Chain of Alcohols in Stabilization of Natural Oil Blends with Diesel

Sample details are:

	Percentage of diesel	Percentage of <i>Jatropha</i> seed oil (SVO)	Percentage of n-butanol	Percentage of t-butanol	Percentage of Iso-butanol
Sample 1	100	—	—	—	—
Sample 2	—	100	—	—	—
Sample 3	—	—	100	—	—
Sample 4	—	—	—	100	—
Sample 5	—	—	—	—	100
Sample 6	95	5	—	—	—
Sample 7	90	10	—	—	—
Sample 8	90	5	5	—	—
Sample 9	85	10	5	—	—
Sample 10	90	5	—	5	—
Sample 11	85	10	—	5	—
Sample 12	90	5	—	—	5
Sample 13	85	10	—	—	5

Results of the experiment are

	0 Days	1 Days	7 Days	30 Days	90 Days	180 Days
Sample 1	No precipitate					
Sample 2	No precipitate					
Sample 3	No precipitate					
Sample 4	No precipitate					
Sample 5	No precipitate					
Sample 6	No precipitate	Precipitate	Precipitate	Precipitate	Precipitate	Precipitate
Sample 7	Precipitate	Precipitate	Precipitate	Precipitate	Precipitate	Precipitate
Sample 8	No precipitate					
Sample 9	No precipitate					
Sample 10	No precipitate	No precipitate	No precipitate	Precipitate	Precipitate	Precipitate
Sample 11	No precipitate	No precipitate	Precipitate	Precipitate	Precipitate	Precipitate
Sample 12	No precipitate	No precipitate	No precipitate	No precipitate	Precipitate	Precipitate
Sample 13	No precipitate	No precipitate	No precipitate	Precipitate	Precipitate	Precipitate

Though iso-butanol and t-butanol has same number of carbons as n-butanol, they fail to stabilize *Jatropha* seed oil (SVO) blend with diesel for more than 30 and 7 days respectively, clearly demonstrating that normal or linear carbon chain of n-butanol played a role. Iso-butanol cannot be used in the composition of present invention as it fails to stabilize blends of *Jatropha* seed oil (SVO) in diesel for at least 30 days.

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The flash points of diesel blends of n-butanol, iso-butanol and tertiary butanol containing 10% *Jatropha* seed oil is as given in the table below.

Sample No	Diesel	n-butanol	Iso-butanol	Tertiary butanol	<i>Jatropha</i> seed oil	Flash point (° C.)
1	100	—	—	—	—	42
2	—	100	—	—	—	37
3	—	—	100	—	—	28
4	—	—	—	100	—	16
5	—	—	—	—	100	>100
6	90	10	—	—	—	37

-continued

Sample No	Diesel	n-butanol	Iso-butanol	Tertiary butanol	<i>Jatropha</i> seed oil	Flash point (° C.)
7	90	—	10	—	—	29
8	90	—	—	10	—	17
9	90	—	—	—	10	52

-continued

Sample No	Diesel	n-butanol	Iso-butanol	Tertiary butanol	Jatropha seed oil	Flash point (° C.)
10	80	10			10	39
11	80		10		10	30
12	80			10	10	17
13	80	20				37.5
14	80		20			29
15	80			20		17
16	80				20	54

Current international norms require diesel to have at least 35° C. as flash point. Observing above results clearly demonstrates that only n-butanol meets the requirement of flash-point for diesel. Iso-butanol and tertiary butanol have low flash point which makes them unfit as blends for diesel and natural oil. Thus it is evident that among the three forms of alcohol, only normal chain alcohol (n-butanol) is fit to stabilize blends of diesel and natural oil.

Performance of a Diesel Engine Fueled by Rapeseed Oil with Normal Chain Alcohols

Sample details are:

Sample number	Diesel	n-butanol	n-pentanol	n-hexanol	Jatropha seed oil
1	100				
2	90	10			
3	90		10		
4	90			10	
5	90				10
6				10	90
7			10		90
8		10			90
9				20	80
10			20		80
11		20			80
12				30	70
13			30		70
14		30			70
15					100

The results are

Sample number	1 day	7 days	14 days	30 days	90 days
1	No precipitate				
2	No precipitate				
3	No precipitate				
4	No precipitate				
5	Precipitate	Precipitate	Precipitate	Precipitate	Precipitate
6	No precipitate				
7	No precipitate	No precipitate	No precipitate	No precipitate	Precipitate
8	No precipitate	No precipitate	No precipitate	No precipitate	Precipitate
9	No precipitate	No precipitate	No precipitate	Precipitate	Precipitate
10	No precipitate	No precipitate	No precipitate	Precipitate	Precipitate
11	No precipitate	No precipitate	No precipitate	Precipitate	Precipitate
12	No precipitate	No precipitate	No precipitate	Precipitate	Precipitate
13	No precipitate	No precipitate	Precipitate	Precipitate	Precipitate
14	No precipitate	No precipitate	Precipitate	Precipitate	Precipitate
15	No precipitate				

The above experiment demonstrates that the blends of Jatropha seed oil (SVO) and normal chain alcohols (n-butanol, n-hexanol, n-pentanol) are not stable in solution for more than 30 days. When alcohol percentage in blend is increased to 20%, the stability decreased to 14 days and when the alcohol percentage is increased to 30%, the stability reduced to one week. This clearly demonstrates that, natural oil blends of normal chain alcohols have less stability and they can be used only in the form of diesel blends of natural oil and normal chain alcohols.

Physico Chemical Analysis of Diesel Blends of Natural Oil in Presence and Absence of Linear Chain Alcohols  
Sample: Commercial Diesel (100% Diesel)

No.	PARAMETERS	RESULTS	Requirements as per IS: 1460: 2000	Requirements as per IS: 1460: 2005	PROTOCOL
1.	Acidity Inorganic	Nil	Nil	Nil	IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	<0.05	0.20 max	To Report	IS: 1448 (P: 2)
3.	Ash, % by wt.,	<0.001	0.01 max	0.01 max	IS: 1448 (P: 4)
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.25	0.30 max	0.3 max	IS: 1448 (P: 8)
5.	Cetane Index	53	46 min	48 min	ASTM D 4737
6.	Pour Point, ° C.	-15	3 max	3 max	IS: 1448 (P-10)
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not worse than No. 1	Not worse than No. 1	IS: 1448 (P-15)
8.			Distillation Recovery, % by vol.		
a.	At 350° C.	90	85 min	85 min	IS: 1448 (P: 18)
b.	At 370° C.	97	95 min	95 min	IS: 1448 (P: 18)
9.	Flash Point, (Abel), ° C.	38	35 min	35 min	IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 40° C., cSt	2.28	2.0-5.0	2.0-5.0	IS: 1448 (P: 25)
11.	Sediments, % by wt.,	0.02	0.05 max	0.05 max	IS: 1448 (P: 30)
12.	Density @ 15° C., Kg/m <sup>3</sup>	818.9	820-860	820-860	IS: 1448 (P: 16)
13.	Total Sulphur, % by wt.,	0.04	0.25 max	0.05 max	IS: 1448 (P: 33)

-continued

No.	PARAMETERS	RESULTS	Requirements	Requirements	PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
14.	Water Content, ppm by wt.,	205	500 max	500 max	IS: 1448 (P: 40)

Sample: 90% Diesel, 10% n-butanol

No.	PARAMETERS	RESULTS	Requirements	Requirements	PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
1.	Acidity Inorganic	Nil	Nil	Nil	IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	0.20	0.20 max	To Report	IS: 1448 (P: 2)
3.	Ash, % by wt.,	<0.001	0.01 max	0.01 max	IS: 1448 (P: 4)
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.3	0.30 max	0.3 max	IS: 1448 (P: 8)
5.	Cetane Index	51	46 min	48 min	ASTM D 4737
6.	Pour Point, ° C.	-12	3 max	3 max	IS: 1448 (P-10)
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not worse than No. 1	Not worse than No. 1	IS: 1448 (P-15)
8.		Distillation Recovery, % by vol.			
A	At 350° C.	85	85 min	85 min	IS: 1448 (P: 18)
B	At 370° C.	96	95 min	95 min	IS: 1448 (P: 18)
9.	Flash Point, (Abel), ° C.	48	35 min	35 min	IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 40° C., cSt	3.22	2.0-5.0	2.0-5.0	IS: 1448 (P: 25)
11.	Sediments, % by wt.,	0.02	0.05 max	0.05 max	IS: 1448 (P: 30)
12.	Density @ 15° C., Kg/m <sup>3</sup>	830.9	820-860	820-860	IS: 1448 (P: 16)
13.	Total Sulphur, % by wt.,	0.04	0.25 max	0.05 max	IS: 1448 (P: 33)
14.	Water Content, ppm by wt.,	178	500 max	500 max	IS: 1448 (P: 40)

Sample: 90% Diesel, 10% Jatropa Oil (Transesterified)

No.	PARAMETERS	RESULTS	Requirements	Requirements	PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
1.	Acidity Inorganic	Nil	Nil	Nil	IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	0.13	0.20 max	To Report	IS: 1448 (P: 2)
3.	Ash, % by wt.,	<0.001	0.01 max	0.01 max	IS: 1448 (P: 4)
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.29	0.30 max	0.3 max	IS: 1448 (P: 8)
5.	Cetane Index	53	46 min	48 min	ASTM D 4737
6.	Pour Point, ° C.	-6	3 max	3 max	IS: 1448 (P-10)
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not worse than No. 1	Not worse than No. 1	IS: 1448 (P-15)
8.		Distillation Recovery, % by vol.			
a	At 350° C.	85	85 min	85 min	IS: 1448 (P: 18)
b	At 370° C.	95	95 min	95 min	IS: 1448 (P: 18)
9.	Flash Point, (Abel), ° C.	42	35 min	35 min	IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 40° C., cSt	2.79	2.0-5.0	2.0-5.0	IS: 1448 (P: 25)
11.	Sediments, % by wt.,	0.03	0.05 max	0.05 max	IS: 1448 (P: 30)
12.	Density @ 15° C., Kg/m <sup>3</sup>	826.4	820-860	820-860	IS: 1448 (P: 16)

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No.	PARAMETERS	RESULTS	Requirements		PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
13.	Total Sulphur, % by wt.,	0.03	0.25 max	0.05 max	IS: 1448 (P: 33)
14.	Water Content, ppm by wt.,	240	500 max	500 max	IS: 1448 (P: 40)

Sample: 90% Diesel, 10% Jatropha Oil (SVO)

No.	PARAMETERS	RESULTS	Requirements		PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
1.	Acidity Inorganic	Nil	Nil	Nil	IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	0.09	0.20 max	To Report	IS: 1448 (P: 2)
3.	Ash, % by wt.,	<0.001	0.01 max	0.01 max	IS: 1448 (P: 4)
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.69	0.30 max	0.3 max	IS: 1448 (P: 8)
5.	Cetane Index	51	46 min	48 min	ASTM D 4737
6.	Pour Point, ° C.	-6	3 max	3 max	IS: 1448 (P-10)
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not worse than No. 1	Not worse than No. 1	IS: 1448 (P-15)
8.		Distillation Recovery, % by vol.			
a	At 350° C.	85	85 min	85 min	IS: 1448 (P: 18)
b	At 370° C.	95	95 min	95 min	IS: 1448 (P: 18)
9.	Flash Point, (Abel), ° C.	48	35 min	35 min	IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 40° C., cSt	2.98	2.0-5.0	2.0-5.0	IS: 1448 (P: 25)
11.	Sediments, % by wt.,	0.08	0.05 max	0.05 max	IS: 1448 (P: 30)
12.	Density @ 15° C., Kg/m <sup>3</sup>	832	820-860	820-860	IS: 1448 (P: 16)
13.	Total Sulphur, % by wt.,	0.12	0.25 max	0.05 max	IS: 1448 (P: 33)
14.	Water Content, ppm by wt.,	272	500 max	500 max	IS: 1448 (P: 40)

Sample: 10% Sunflower Oil (SVO), 90% Diesel

S. No.	PARAMETERS	RESULTS	Requirements		PROTOCOL
			as per IS: 1460: 2005		
1.	Acidity Inorganic	Nil	Nil		IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	0.01	To Report		IS: 1448 (P: 2)
3.	Ash, % by wt.,	0.001	0.01 max		IS: 1448 (P: 4)
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.54	0.3 max		IS: 1448 (P: 8)
5.	Cetane Index	57	48 min		ASTM D 976
6.	Pour Point, ° c.	-12	3 max		IS: 1448 (P-10)
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not Worse than No. 1		IS: 1448 (P-15)
8.		Distillation Recovery, Temp U <sub>c</sub>			
a	at 350° C.	86	85 min		IS: 1448 (P: 18)
b	at 370° C.	97	95 min		IS: 1448 (P: 18)
9.	Flash Point (Abel), ° c.	50	35 min		IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 40° C. cSt	2.84	2.0-5.0		IS: 1448 (P: 25)
11.	Sediments, % by wt.,	0.026	0.05 max		IS: 1448 (P: 37)
12.	Density @ 15° C. g/ml	0.8264	820-860		IS: 1448 (P: 16)
13.	Total Sulphur, % by wt.,	0.14	0.05 max		IS: 1448 (P: 33)
14.	Water Content, ppm by wt.,	179	500 max		IS: 1448 (P: 40)

Sample: 10% Neem Seed Oil (SVO), 90% Diesel

S. No.	PARAMETERS	RESULTS	Requirements		PROTOCOL
			as per IS: 1460: 2005		
1.	Acidity Inorganic	Nil	Nil		IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	0.85	To Report		IS: 1448 (P: 2)
3.	Ash, % by wt.,	0.001	0.01 max		IS: 1448 (P: 4)
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.96	0.3 max		IS: 1448 (P: 8)
5.	Cetane Index	53	48 min		ASTM D 976
6.	Pour Point ° C.	-18	3 max		IS: 1448 (P-10)
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not Worse than No. 1		IS: 1448 (P-15)
8.		Distillation Recovery, Temp ° C.			
a	at 350° C.	88	85 min		IS: 1448 (P: 18)
b	at 370° C.	96	95 min		IS: 1448 (P: 18)
9.	Flash Point (Abel) ° C.	44	35 min		IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 40° C. cSt	2.04	2.0-5.0		IS: 1448 (P: 25)
11.	Sediments, % by wt.,	0.012	0.05 max		IS: 1448 (P: 37)
12.	Density @ 15° C. Q/ml	0.8308	820-860		IS: 1448 (P: 16)
13.	Total Sulphur, % by wt.,	0.14	0.05 max		IS: 1448 (P: 33)
14.	Water Content, ppm by wt.,	259	500 max		IS: 1448 (P: 40)

Sample: 85% Diesel, 7.5% Jatropha Oil, 7.5% n-butanol

No.	PARAMETERS	RESULTS	Requirements		PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
1.	Acidity Inorganic	Nil	Nil	Nil	IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	0.12	0.20 max	To Report	IS: 1448 (P: 2)
3.	Ash, % by wt.,	<0.001	0.01 max	0.01 max	IS: 1448 (P: 4)
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.29	0.30 max	0.3 max	IS: 1448 (P: 8)
5.	Cetane Index	52	46 min	48 min	ASTM D 4737
6.	Pour Point, ° C.	-12	3 max	3 max	IS: 1448 (P-10)
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not worse than No. 1	Not worse than No. 1	IS: 1448 (P-15)
8.		Distillation Recovery, % by vol.			
A	At 350° C.	85	85 min	85 min	IS: 1448 (P: 18)
B	At 370° C.	96	95 min	95 min	IS: 1448 (P: 18)
9.	Flash Point, (Abel), ° C.	48	35 min	35 min	IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 40° C., cSt	3.61	2.0-5.0	2.0-5.0	IS: 1448 (P: 25)
11.	Sediments, % by wt.,	0.03	0.05 max	0.05 max	IS: 1448 (P: 30)
12.	Density @ 15° C., Kg/m <sup>3</sup>	836.9	820-860	820-860	IS: 1448 (P: 16)
13.	Total Sulphur, % by wt.,	0.04	0.25 max	0.05 max	IS: 1448 (P: 33)
14.	Water Content, ppm by wt.,	239	500 max	500 max	IS: 1448 (P: 40)

Sample: 80% Diesel, 10% Jatropha Oil, 10% n-butanol

No.	PARAMETERS	RESULTS	Requirements		PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
1.	Acidity Inorganic	Nil	Nil	Nil	IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	0.1	0.20 max	To Report	IS: 1448 (P: 2)
3.	Ash, % by wt.,	<0.001	0.01 max	0.01 max	IS: 1448 (P: 4)

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No.	PARAMETERS	RESULTS	Requirements		PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.29	0.30 max	0.3 max	IS: 1448 (P: 8)
5.	Cetane Index	50	46 min	48 min	ASTM D 4737
6.	Pour Point, ° C.	-12	3 max	3 max	IS: 1448 (P-10)
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not worse than No. 1	Not worse than No. 1	IS: 1448 (P-15)
8.		Distillation Recovery, % by vol.			
A	At 350° C.	85	85 min	85 min	IS: 1448 (P: 18)
B	At 370° C.	96	95 min	95 min	IS: 1448 (P: 18)
9.	Flash Point, (Abel), ° C.	46	35 min	35 min	IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 40° C., cSt	4.29	2.0-5.0	2.0-5.0	IS: 1448 (P: 25)
11.	Sediments, % by wt.,	0.03	0.05 max	0.05 max	IS: 1448 (P: 30)
12.	Density @ 15° C., Kg/m <sup>3</sup>	841.9	820-860	820-860	IS: 1448 (P: 16)
13.	Total Sulphur, % by wt.,	0.04	0.25 max	0.05 max	IS: 1448 (P: 33)
14.	Water Content, ppm by wt.,	237	500 max	500 max	IS: 1448 (P: 40)

Sample: 85% Diesel, 7.5% Neem oil, 7.5% n-butanol

No.	PARAMETERS	RESULTS	Requirements		PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
1.	Acidity Inorganic	Nil	Nil	Nil	IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	0.08	0.20 max	To Report	IS: 1448 (P: 2)
3.	Ash, % by wt.,	<0.001	0.01 max	0.01 max	IS: 1448 (P: 4)
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.27	0.30 max	0.3 max	IS: 1448 (P: 8)
5.	Cetane Index	51	46 min	48 min	ASTM D 4737
6.	Pour Point, ° C.	-11	3 max	3 max	IS: 1448 (P-10)
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not worse than No. 1	Not worse than No. 1	IS: 1448 (P-15)
8.		Distillation Recovery, % by vol.			
A	At 350° C.	87	85 min	85 min	IS: 1448 (P: 18)
B	At 370° C.	97	95 min	95 min	IS: 1448 (P: 18)
9.	Flash Point, (Abel), ° C.	39	35 min	35 min	IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 40° C., cSt	3.11	2.0-5.0	2.0-5.0	IS: 1448 (P: 25)
11.	Sediments, % by wt.,	0.02	0.05 max	0.05 max	IS: 1448 (P: 30)
12.	Density @ 15° C., Kg/m <sup>3</sup>	829.7	820-860	820-860	IS: 1448 (P: 16)
13.	Total Sulphur, % by wt.,	0.05	0.25 max	0.05 max	IS: 1448 (P: 33)
14.	Water Content, ppm by wt.,	228	500 max	500 max	IS: 1448 (P: 40)

Sample: 80% Diesel, 10% Neem Oil, 10% n-butanol

No.	PARAMETERS	RESULTS	Requirements		PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
1.	Acidity Inorganic	Nil	Nil	Nil	IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	0.12	0.20 max	To Report	IS: 1448 (P: 2)
3.	Ash, % by wt.,	<0.001	0.01 max	0.01 max	IS: 1448 (P: 4)
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.29	0.30 max	0.3 max	IS: 1448 (P: 8)
5.	Cetane Index	50	46 min	48 min	ASTM D 4737
6.	Pour Point, ° C.	-11	3 max	3 max	IS: 1448 (P-10)

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No.	PARAMETERS	RESULTS	Requirements		PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not worse than No. 1	Not worse than No. 1	IS: 1448 (P-15)
8.			Distillation Recovery, % by vol.		
A	At 350° C.	86	85 min	85 min	IS: 1448 (P: 18)
B	At 370° C.	95	95 min	95 min	IS: 1448 (P: 18)
9.	Flash Point, (Abel), ° C.	38	35 min	35 min	IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 40° C., cSt	3.78	2.0-5.0	2.0-5.0	IS: 1448 (P: 25)
11.	Sediments, % by wt.,	0.03	0.05 max	0.05 max	IS: 1448 (P: 30)
12.	Density @ 15° C., Kg/m <sup>3</sup>	835.7	820-860	820-860	IS: 1448 (P: 16)
13.	Total Sulphur, % by wt.,	0.05	0.25 max	0.05 max	IS: 1448 (P: 33)
14.	Water Content, ppm by wt.,	273	500 max	500 max	IS: 1448 (P: 40)

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Sample: 90% Diesel, 5% Neem Oil, 5% n-butanol

No.	PARAMETERS	RESULTS	Requirements		PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
1.	Acidity Inorganic	Nil	Nil	Nil	IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	<0.05	0.20 max	To Report	IS: 1448 (P: 2)
3.	Ash, % by wt.,	<0.001	0.01 max	0.01 max	IS: 1448 (P: 4)
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.26	0.30 max	0.3 max	IS: 1448 (P: 8)
5.	Cetane Index	52	46 min	48 min	ASTM D 4737
6.	Pour Point, ° C.	-11	3 max	3 max	IS: 1448 (P-10)
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not worse than No. 1	Not worse than No. 1	IS: 1448 (P-15)
8.			Distillation Recovery, % by vol.		
a	At 350° C.	90	85 min	85 min	IS: 1448 (P: 18)
b	At 370° C.	97	95 min	95 min	IS: 1448 (P: 18)
9.	Flash Point, (Abel), ° C.	37	35 min	35 min	IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 40° C., cSt	2.81	2.0-5.0	2.0-5.0	IS: 1448 (P: 25)
11.	Sediments, % by wt.,	0.04	0.05 max	0.05 max	IS: 1448 (P: 30)
12.	Density @ 15° C., Kg/m <sup>3</sup>	827.3	820-860	820-860	IS: 1448 (P: 16)
13.	Total Sulphur, % by wt.,	0.04	0.25 max	0.05 max	IS: 1448 (P: 33)
14.	Water Content, ppm by wt.,	198	500 max	500 max	IS: 1448 (P: 40)

Sample: 85% Diesel, 7.5% Sunflower Oil, 7.5% n-butanol

No.	PARAMETERS	RESULTS	Requirements		PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
1.	Acidity Inorganic	Nil	Nil	Nil	IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	<0.05	0.20 max	To Report	IS: 1448 (P: 2)
3.	Ash, % by wt.,	<0.001	0.01 max	0.01 max	IS: 1448 (P: 4)
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.29	0.30 max	0.3 max	IS: 1448 (P: 8)
5.	Cetane Index	53	46 min	48 min	ASTM D 4737
6.	Pour Point, ° C.	-10	3 max	3 max	IS: 1448 (P-10)
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not worse than No. 1	Not worse than No. 1	IS: 1448 (P-15)

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No.	PARAMETERS	RESULTS	Requirements		PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
8.	Distillation Recovery, % by vol.				
a	At 350° C.	88	85 min	85 min	IS: 1448 (P: 18)
b	At 370° C.	96	95 min	95 min	IS: 1448 (P: 18)
9.	Flash Point, (Abel), ° C.	37	35 min	35 min	IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 40° C., cSt	2.92	2.0-5.0	2.0-5.0	IS: 1448 (P: 25)
11.	Sediments, % by wt.,	0.05	0.05 max	0.05 max	IS: 1448 (P: 30)
12.	Density @ 15° C., Kg/m <sup>3</sup>	832	820-860	820-860	IS: 1448 (P: 16)
13.	Total Sulphur, % by wt.,	0.05	0.25 max	0.05 max	IS: 1448 (P: 33)
14.	Water Content, ppm by wt.,	175	500 max	500 max	IS: 1448 (P: 40)

Sample: 90% Diesel, 5% Sunflower Oil, 5% n-butanol

No.	PARAMETERS	RESULTS	Requirements		PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
1.	Acidity Inorganic	Nil	Nil	Nil	IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	<0.05	0.20 max	To Report	IS: 1448 (P: 2)
3.	Ash, % by wt.,	<0.001	0.01 max	0.01 max	IS: 1448 (P: 4)
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.24	0.30 max	0.3 max	IS: 1448 (P: 8)
5.	Cetane Index	52	46 min	48 min	ASTM D 4737
6.	Pour Point, ° C.	-10	3 max	3 max	IS: 1448 (P-10)
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not worse than No. 1	Not worse than No. 1	IS: 1448 (P-15)
8.	Distillation Recovery, % by vol.				
a	At 350° C.	94	85 min	85 min	IS: 1448 (P: 18)
b	At 370° C.	97	95 min	95 min	IS: 1448 (P: 18)
9.	Flash Point, (Abel), ° C.	37	35 min	35 min	IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 40° C., cSt	2.61	2.0-5.0	2.0-5.0	IS: 1448 (P: 25)
11.	Sediments, % by wt.,	0.01	0.05 max	0.05 max	IS: 1448 (P: 30)
12.	Density @ 15° C., Kg/m <sup>3</sup>	820	820-860	820-860	IS: 1448 (P: 16)
13.	Total Sulphur, % by wt.,	0.03	0.25 max	0.05 max	IS: 1448 (P: 33)
14.	Water Content, ppm by wt.,	156	500 max	500 max	IS: 1448 (P: 40)

Sample: 80% Diesel, 10% Sunflower Oil, 10% n-butanol

No.	PARAMETERS	RESULTS	Requirements		PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
1.	Acidity Inorganic	Nil	Nil	Nil	IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	<0.05	0.20 max	To Report	IS: 1448 (P: 2)
3.	Ash, % by wt.,	<0.001	0.01 max	0.01 max	IS: 1448 (P: 4)
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.30	0.30 max	0.3 max	IS: 1448 (P: 8)
5.	Cetane Index	53	46 min	48 min	ASTM D 4737
6.	Pour Point, ° C.	-11	3 max	3 max	IS: 1448 (P-10)
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not worse than No. 1	Not worse than No. 1	IS: 1448 (P-15)
8.	Distillation Recovery, % by vol.				
a	At 350° C.	86	85 min	85 min	IS: 1448 (P: 18)
b	At 370° C.	96	95 min	95 min	IS: 1448 (P: 18)

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No.	PARAMETERS	RESULTS	Requirements	Requirements	PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
9.	Flash Point, (Abel), ° C.	38	35 min	35 min	IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 400 C., cSt	3.02	2.0-5.0	2.0-5.0	IS: 1448 (P: 25)
11.	Sediments, % by wt.,	0.05	0.05 max	0.05 max	IS: 1448 (P: 30)
12.	Density @ 15° C., Kg/m <sup>3</sup>	843	820-860	820-860	IS: 1448 (P: 16)
13.	Total Sulphur, % by wt.,	0.05	0.25 max	0.05 max	IS: 1448 (P: 33)
14.	Water Content, ppm by wt.,	205	500 max	500 max	IS: 1448 (P: 40)

Sample: 80% Diesel, 10% Jatropha Oil, 10% Cardanol

No.	PARAMETERS	RESULTS	Requirements	Requirements	PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
1.	Acidity Inorganic	Nil	Nil	Nil	IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	<0.05	0.20 max	To Report	IS: 1448 (P: 2)
3.	Ash, % by wt.,	<0.001	0.01 max	0.01 max	IS: 1448 (P: 4)
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.27	0.30 max	0.3 max	IS: 1448 (P: 8)
5.	Cetane Index	53	46 min	48 min	ASTM D 4737
6.	Pour Point, ° C.	-12	3 max	3 max	IS: 1448 (P-10)
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not worse than No. 1	Not worse than No. 1	IS: 1448 (P-15)
8.	Distillation Recovery, % by vol.				
a	At 350° C.	90	85 min	85 min	IS: 1448 (P: 18)
b	At 370° C.	97	95 min	95 min	IS: 1448 (P: 18)
9.	Flash Point, (Abel), ° C.	46	35 min	35 min	IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 40° C., cSt	2.71	2.0-5.0	2.0-5.0	IS: 1448 (P: 25)
11.	Sediments, % by wt.,	0.03	0.05 max	0.05 max	IS: 1448 (P: 30)
12.	Density @15° C., Kg/m <sup>3</sup>	827.9	820-860	820-860	IS: 1448 (P: 16)
13.	Total Sulphur, % by wt.,	0.01	0.25 max	0.05 max	IS: 1448 (P: 33)
14.	Water Content, ppm by wt.,	178	500 max	500 max	IS: 1448 (P: 40)

Sample: 85% Diesel, 7.5% Jatropha Oil, 7.5% Cardanol

No.	PARAMETERS	RESULTS	Requirements	Requirements	PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
1.	Acidity Inorganic	Nil	Nil	Nil	IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	<0.05	0.20 max	To Report	IS: 1448 (P: 2)
3.	Ash, % by wt.,	<0.001	0.01 max	0.01 max	IS: 1448 (P: 4)
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.29	0.30 max	0.3 max	IS: 1448 (P: 8)
5.	Cetane Index	52	46 min	48 min	ASTM D 4737
6.	Pour Point, ° C.	-9	3 max	3 max	IS: 1448 (P-10)
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not worse than No. 1	Not worse than No. 1	IS: 1448 (P-15)
8.	Distillation Recovery, % by vol.				
a	At 350° C.	88	85 min	85 min	IS: 1448 (P: 18)
b	At 370° C.	96	95 min	95 min	IS: 1448 (P: 18)
9.	Flash Point, (Abel), ° C.	50	35 min	35 min	IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 40° C., cSt	2.89	2.0-5.0	2.0-5.0	IS: 1448 (P: 25)

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No.	PARAMETERS	RESULTS	Requirements	Requirements	PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
11.	Sediments, % by wt.,	0.03	0.05 max	0.05 max	IS: 1448 (P: 30)
12.	Density @ 15° C., Kg/m <sup>3</sup>	833.9	820-860	820-860	IS: 1448 (P: 16)
13.	Total Sulphur, % by wt.,	0.05	0.25 max	0.05 max	IS: 1448 (P: 33)
14.	Water Content, ppm by wt.,	185	500 max	500 max	IS: 1448 (P: 40)

Sample: 90% Diesel, 5% Jatropa Oil, 5% n-butanol

No.	PARAMETERS	RESULTS	Requirements	Requirements	PROTOCOL
			as per IS: 1460: 2000	as per IS: 1460: 2005	
1.	Acidity Inorganic	Nil	Nil	Nil	IS: 1448 (P: 2)
2.	Total Acidity, mg KOH/g	<0.05	0.20 max	To Report	IS: 1448 (P: 2)
3.	Ash, % by wt.,	<0.001	0.01 max	0.01 max	IS: 1448 (P: 4)
4.	Carbon Residue Ramsbottom, on 10% residue by wt.,	0.26	0.30 max	0.3 max	IS: 1448 (P: 8)
5.	Cetane Index	52	46 min	48 min	ASTM D 4737
6.	Pour Point, ° C.	-9	3 max	3 max	IS: 1448 (P-10)
7.	Copper Strip Corrosion for 3 Hrs @ 100° C.	1a	Not worse than No. 1	Not worse than No. 1	IS: 1448 (P-15)
8.		Distillation Recovery, % by vol.			
a	At 350° C.	94	85 min	85 min	IS: 1448 (P: 18)
b	At 370° C.	97	95 min	95 min	IS: 1448 (P: 18)
9.	Flash Point, (Abel), ° C.	48	35 min	35 min	IS: 1448 (P: 20)
10.	Kinematic Viscosity @ 40° C., cSt	2.62	2.0-5.0	2.0-5.0	IS: 1448 (P: 25)
11.	Sediments, % by wt.,	0.01	0.05 max	0.05 max	IS: 1448 (P: 30)
12.	Density @15° C., Kg/m <sup>3</sup>	820.9	820-860	820-860	IS: 1448 (P: 16)
13.	Total Sulphur, % by wt.,	0.04	0.25 max	0.05 max	IS: 1448 (P: 33)
14.	Water Content, ppm by wt.,	178	500 max	500 max	IS: 1448 (P: 40)

The physico chemical analysis was done as per IS: 1460: 2000 and IS: 1460: 2005, the Indian standard for euro diesel. The analysis was done immediately after preparing the samples. As evident from the above data, the physico chemical characteristics of the blends of n-butanol with natural oils (Sunflower oil, Jatropa oil, neem seed oil) and diesel satisfy the parameters of IS: 1460: 2000 and IS: 1460: 2005, while the combination of natural oil and diesel does not.

Engine Testing of Natural Oil as Blend of Diesel—Effect of n-butanol and Cardanol on Engine Performance and Emission

Engine Specification:

Make: Kirloskar oil engines

No of cylinders: twin cylinder

Bore: 102 mm

Stroke: 116 mm

Displacement: 1.896 liters

Compression ratio: 17.5

Bhp: 20 hp

Injection pressure: 185 bar

Loading Sequence Followed:

a) No load (0)

b) 3.1 kW

c) 6.1 kW

d) 9.1 kW

e) 11.1 kW

Pressure sensor: Kistler (maximum pressure: 250 bar)  
ASTM standards are followed to measure the air flow using  
an air drum with a calibrated orifice and a manometer  
Measured room temperature: 30±3° C.

Air density: 1.164 Kg/m<sup>3</sup>

Test Parameters:

mf/min: Grams of fuel consumed per minute

Load kW: Load applied for specific study

Rpm: Engine rotations per minute

HC: Total hydrocarbons produced in exhaust smoke

CO: Total carbon monoxide produced in exhaust smoke

CO<sub>2</sub>: Total carbon dioxide produced in exhaust smoke

For each fuel blend, three separate experiments (sets 1-3) were done and average was taken. Engine was off for atleast one hour between experiments involving same fuel and three hours when engine was switched to a new fuel. For every experiment, engine was run for atleast 30 mins to get rid of any residual fuel in fuel lines before recording parameters.

The tested samples are

Commercial diesel

Sample 1: 90% Diesel, 5% Jatropa oil (SVO), 5% n-butanol

Sample 2: 85% Diesel, 7.5% Jatropa oil (SVO), 7.5% n-butanol

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- Sample 3: 80% Diesel, 10% Jatropa oil (SVO), 10% n-butanol
- Sample 4: 95% Diesel, 5% n-butanol
- Sample 5: 90% Diesel, 10% n-butanol
- Sample 6: 90% Diesel, 5% Jatropa oil (SVO), 5% Cardanol
- Sample 7: 85% Diesel, 7.5% Jatropa oil (SVO), 7.5% Cardanol

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- Sample 8: 80% Diesel, 10% Jatropa oil (SVO), 10% Cardanol
  - Sample 9: 95% Diesel, 5% Cardanol
  - Sample 10: 90% Diesel, 10% Jatropa oil (SVO)
  - Sample 11: 90% Diesel, 10% Jatropa biodiesel (transesterified)
  - Sample 12: 85% Diesel, 15% Jatropa biodiesel (transesterified)
- Hydrocarbon Emissions from Exhaust for Above Samples

Hydrocarbon vs Load							
Load	Diesel	Sample 1	Sample 2	Sample 3	Sample 4	Sample 5	Sample 6
0	24.0	24	30.0	34.0	31.5	27.0	21.0
3	7.5	8	10.0	11.0	14.5	22.0	17.0
6.1	10.0	10.0	12.0	15.0	20.5	23.0	14.5
9	20.0	16.5	18.0	19.5	21.5	25.5	26.0
11.1	31.0	27.5	31	32	35.5	38.5	43.0

Load	Sample 7	Sample 8	Sample 9	Sample 10	Sample 11	Sample 12
0	21.0	16.0	35.	40.0	26.5	32.0
3	13.5	15.0	18.	20.0	17.0	19.5
6.1	14.5	15.5	20.0	23.0	14.0	15.5
9	24.5	32.0	35.0	38.0	18.0	22.0
11.1	36.5	40.0	43.0	49.0	34.5	38.0

Carbon Monoxide Emissions from Above Samples

Carbon monoxide vs Load							
Load	Diesel	Sample 1	Sample 2	Sample 3	Sample 4	Sample 5	Sample 6
0	0.06	0.02	0.08	0.08	0.05	0.03	0.02
3	0.07	0.06	0.05	0.07	0.06	0.06	0.05
6.1	0.07	0.07	0.07	0.07	0.08	0.07	0.08
9	0.10	0.11	0.13	0.11	0.11	0.09	0.12
11.1	0.30	0.23	0.23	0.32	0.27	0.25	0.35

Load	Sample 7	Sample 8	Sample 9	Sample 10	Sample 11	Sample 12
0	0.02	0.06	0.09	0.10	0.07	0.07
3	0.07	0.10	0.12	0.09	0.09	0.08
6.1	0.08	0.13	0.17	0.20	0.09	0.10
9	0.13	0.18	0.24	0.35	0.13	0.12
11.1	0.43	0.45	0.47	0.56	0.30	0.36

Carbon Dioxide Emissions for Above Samples

Carbon dioxide vs Load							
Load	Diesel	Sample 1	Sample 2	Sample 3	Sample 4	Sample 5	Sample 6
0	2.6	2.1	2.3	2.2	2.6	2.0	2.2
3	4.5	4.6	4.4	4.4	4.7	4.5	4.9
6.1	6.2	6.4	6.3	6.3	6.2	6.3	6.6
9	8.5	8.6	8.4	8.5	8.5	8.6	8.6
11.1	10.3	10.4	10.2	10.3	10.3	10.2	10.4

Load	Sample 7	Sample 8	Sample 9	Sample 10	Sample 11	Sample 12
0	2.1	2.7	2.9	3.1	2.8	2.2
3	4.6	4.6	4.9	5.6	4.5	4.4
6.1	6.5	6.4	6.6	7.5	6.4	6.3
9	8.7	8.9	8.7	10.6	8.6	8.5
11.1	10.5	10.6	10.4	12.4	10.1	10.1

Fuel Consumption for Above Samples

Fuel consumption Vs Load							
Load	Diesel	Sample 1	Sample 2	Sample 3	Sample 4	Sample 5	Sample 6
0.0	18.0	18.0	18.0	17.7	18.8	16.8	18.0
3.0	26.7	27.0	26.3	25.8	26.3	25.8	27.0
6.1	36.0	35.7	35.7	36.2	35.8	35.0	36.0
9.0	47.0	45.7	47.0	46.0	45.8	46.0	45.0
11.1	56.7	56.0	56.2	56.2	55.2	54.5	55.0

Load	Sample 7	Sample 8	Sample 9	Sample 10	Sample 11	Sample 12
0.0	16.8	16.7	19.0	19.8	18.0	18.0
3.0	25.8	26.0	27.8	29.7	26.3	26.0
6.1	35.2	36.0	37.8	39.7	35.7	35.8
9.0	45.7	46.5	49.0	49.7	46.0	46.0
11.1	56.0	57.3	58.8	59.7	55.0	56.0

Samples 1-3 (SVO-diesel blends with n-butanol), 6-8 (SVO-diesel blends with cardanol) have fuel parameters (both performance & emission) similar to Sample 11-12 (Biodiesel blends) and commercial diesel. This indicates the proposed normal chain alcohols enable the natural oil blends of diesel to behave similar to commercial diesel.

Engine Test—To Compare Opacity of Different Natural Oil Blends in Diesel

The separation of layers in natural oil-diesel blend cause improper spraying of blend, which results in increased smoke and addition of normal chain alcohols like n-butanol addresses the problem.

When a poor/uneven/non-uniform diesel or diesel blend is injected to diesel engine, due to formation of carbon black deposits on injector nozzle, there could be uneven spraying of fuel, which results in large amounts of smoke. Hence, in a uniform running engine, under standard operation conditions, sudden observance of large amounts of smoke can be attributed to uneven spraying of fuel which in turn can be used to identify faulty/bad fuel.

Experiment design: Diesel engine was connected with container having 10 liters of diesel and ran till it stopped. This was continued for three times and observed any awkward smoke increase. It was observed that engine on average run for 5 hours under a load of 6.1 kW. During last three minutes before stoppage, engine gave an up to 3 fold increase in smoke for all fuel blends and diesel. Similarly, when engine was cranked, first few seconds resulted in lot of smoke. Hence, it was concluded that just after cranking and just before stopping, engine tends to give more smoke. It was also observed that, even though engine was made to run at constant speed, due to constant running, there is a variation of up to 20% in smoke from the exhaust.

Test Conditions

Engine: A Kirloskar twin cylinder with 102 mm bore, 116 mm stroke, 1.9 liters displacement, with compression ratio of 17.5 and 20 hp diesel engine was used.

Total fuel added to storage tank: 10 liters of well mixed diesel

Total runtime: Approximately 5 hours

Engine speed (rpm): 1530±10, system tries to maintain constant speed by adjusting the fuel flow

Opacity monitored window: 30 minutes after cranking engine and 30 minutes before engine self-stoppage (about 4 hours for each fuel blend and diesel).

Tolerable change in opacity: up to 50% change from average reading

Temperature of study: 30±5° C.

Number of tests per fuel/fuel blend: Three tests and average result is taken

Opacity: Usually after 15 minutes of cranking the engine, most of the residual fuel will be burnt and the opacity will be of the fuel being tested. Three readings were taken from the opacity between 20 and 30 minutes and average of that value is taken as the average opacity (starting value).

Sudden stoppage of engine: Sudden stoppage of engine on its own after 30 minutes equilibration (for next four hours) is monitored. In case of an engine stoppage, engine was restarted and five minutes equilibration window is provided before the parameters were recorded again.

It is important to note that the average opacity of specific fuel is compared with its number of times awkward changes in opacity (>50% of average value).

Sample Details:

Diesel: Bharat stage-III diesel from a commercial depot in Bangalore

Sample 1: 20% trans esterified (methyl) jatropha biodiesel, 80% of diesel

Sample 2: 10% trans esterified (methyl) jatropha biodiesel, 90% diesel

Sample 3: 20% Jatropha oil (SVO), 80% diesel

Sample 4: 10% Jatropha oil (SVO), 90% diesel

Sample 5: 5% Jatropha oil, 5% n-butanol, 90% diesel

Sample 6: 10% Jatropha oil, 10% n-butanol, 80% diesel

Sample 7: 5% Jatropha oil, 5% cardanol, 90% diesel

Sample 8: 10% Jatropha oil, 10% cardanol, 80% diesel

Between each experiment engine was off for minimum of 3 hours to let it cool.

Every time a fuel is run, fuel injector system was dismantled and cleaned (washing with diesel, opening nozzle and removing carbon and then checking whether nozzle moving freely, using copper brush remove carbon deposits, etc).

Fuel/Fuel blend	Engine run time (min)			Opacity (%)				No of times spike (>50% increase)			No of times sudden stoppage of engine					
	Average			Average				Average			Average					
Diesel	312	324	317	317.7	21	22	23	22	0	1	1	0.7	0	0	0	0
Sample 1	306	309	301	305.3	29	32	31	30.7	1	2	1	1.3	0	0	0	0

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Fuel/Fuel blend	Engine run time (min)				Opacity (%)				No of times spike (>50% increase)			No of times sudden stoppage of engine				
	Average				Average				Average			Average				
Sample 2	311	306	322	313	20	22	22	21.3	1	1	0	0.7	0	0	0	0
Sample 3	298	292	301	297	37	35	36	36	15	12	13	13.3	2	3	2	2.3
Sample 4	287	289	291	289	31	29	28	29.3	7	8	9	8	1	1	1	1
Sample 5	292	319	309	312.3	14	15	15	16.7	0	1	1	0.7	0	0	0	0
Sample 6	309	327	322	319.3	19	17	19	18.3	1	2	1	1.3	0	0	0	0
Sample 7	312	317	315	314.7	21	22	21	21.3	0	1	1	0.7	0	0	0	0
Sample 8	329	333	317	326.3	27	29	29	28.3	1	1	1	1	0	0	0	0

It is clear from the above table that:

1. Total engine run times of samples containing blends of natural oil and diesel are lower. Addition of n-butanol to the blends of natural oil and diesel seem to address the problem.
2. Number of times engine stopped suddenly is higher for samples containing blends of natural oil and diesel. Addition of n-butanol to the blends of natural oil and diesel seem to address the problem.
3. Number of times spike or sudden increase in opacity observed was more for samples containing blends of natural oil and diesel. Addition of n-butanol to the blends of natural oil and diesel seem to address the problem.
4. Thus, it clearly demonstrates that there is improper running of engine when natural oil and diesel blends are used and addition of n-butanol (normal chain alcohols) to the blends of natural oil and diesel address this problem.

Decrease in NOx Due to Addition of n-butanol and n-hexanol

Engine: A Kirloskar twin cylinder with 102 mm bore, 116 mm stroke, 1.9 liters displacement, with compression ratio of 17.5 and 20 hp diesel engine was used.

SI	mf/min	Load kW	rpm	Nox	Opacity	Egt
Diesel						
1	18	0	1588	135	2	187
1	18	0	1588	135	2	187
2	27	3.1	1551	430	10	242
3	36	6.1	1535	852	19	318
4	47	9	1506	1089	42	419
5	57	11.1	1490	1323	67	490
10% <i>Jatropha</i> SVO, 90% diesel						
1	20	0	1558	384	2	198
2	29	3.1	1556	650	13	257
3	38	6.1	1538	917	29	340
4	49	9.1	1513	1179	54	448
5	58	11.1	1496	1449	79	502
10% <i>Jatropha</i> SVO, 10% n-butanol, 80% diesel						
1	18	0	1562	221	2	185
2	27	3.1	1550	427	8	221
3	36	6	1531	833	19	284
4	46	9	1510	1000	35	409
5	56	11	1497	1080	60	491
10% <i>Jatropha</i> trans esterified biodiesel, 90% diesel						
1	19	0	1576	147	1	183
2	28	3.1	1548	447	10	251
3	39	6	1533	711	19	339
4	48	9	1507	999	42	437
5	59	11.1	1492	1381	66	506

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SI	mf/min	Load kW	rpm	Nox	Opacity	Egt
10% <i>Jatropha</i> SVO, 10% n-hexanol, 80% diesel						
1	18	0	1547	235	1	179
2	27	3.1	1545	410	16	236
3	36	6	1537	810	38	310
4	48	9	1511	1022	50	402
5	57	11	1496	1171	69	489
10% <i>Jatropha</i> SVO, 10% cardanol, 80% diesel						
1	18	0	1556	300	0	183
2	27	3	1545	617	6.9	240
3	38	6	1533	900	20	287
4	49	9	1515	1112	43	401
5	59	11	1496	1184	65	475
Above study clearly demonstrates that NOx emissions from blends of natural oil and diesel are higher compared to diesel alone and they are reduced by addition of n-butanol, n-hexanol and cardanol.						
Decrease in NOx Due to Addition of n-dodecanol, n-decanol, 9Z, 12Z-octadecadien-1-ol						
Engine: A Kirloskar twin cylinder with 102 mm bore, 116 mm stroke, 1.9 liters displacement, with compression ratio of 17.5 and 20 hp diesel engine was used.						
Diesel						
18	0	1554	292	2	185	
27	3	1540	602	7	243	
38	6	1537	904	21	304	
49	9	1517	1135	47	408	
59	11	1499	1394	69	490	
15% <i>Jatropha</i> SVO, 85% diesel						
20	0	1555	391	2	196	
30	3.1	1545	748	13	252	
39	6	1531	1045	30	336	
49	9.1	1512	1269	54	436	
59	11.1	1333	1480	77	500	
7.5% <i>Jatropha</i> SVO, 7.5% n-decanol, 85% diesel						
18	0	1554	315	1	189	
28	3.1	1540	639	3	256	
38	6	1532	876	14	332	
48	9.1	1509	1074	37	434	
58	11	1496	1129	69	510	
15% <i>Jatropha</i> trans esterified biodiesel, 85% diesel						
19	0	1553	363	2	196	
29	3	1535	681	8	268	
38	6.1	1526	898	14	334	
49	9	1503	1115	43	430	
59	11	1492	1331	69	522	





























































-continued

Sample no	0 Days	1 Days	7 Days	14 Days	30 Days	90 Days	120 Days
13	No precipitate						
14	No precipitate						
15	No precipitate						
16	No precipitate						
17	No precipitate						
18	No precipitate						

We claim:

1. A biofuel composition consisting of normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration ranging from about 1% to about 24% v/v, natural oil(s) at a concentration ranging from about 1% to about 24% v/v and diesel at a concentration ranging from about 52% to about 98% v/v, optionally along with fuel additive(s), wherein the natural oil(s) comprise triglycerides.

2. The biofuel composition as claimed in claim 1, wherein the normal chain alcohols are selected from the group consisting of n-butanol, n-pentanol, n-hexanol, n-heptanol, n-octanol (capryl alcohol), n-nonanol (pelargonic alcohol), n-decanol (capric alcohol), n-dodecanol (lauryl alcohol), n-pentadecanol, n-hexadecanol (cetyl alcohol), n-tetradecanol (myristyl alcohol), cis-9-hexadecen-1-ol (palmitoleyl alcohol), n-octadecanol (stearyl alcohol), 9E-octadecen-1-ol (elaidyl alcohol), cis-9-octadecen-1-ol (oleyl alcohol), 9Z, 12Z-octadecadien-1-ol (linoleyl alcohol), 9E, 12E-octadecadien-1-ol (elaidolinoleyl alcohol), 9Z, 12Z, 15Z-octadecatrien-1-ol (linolenyl alcohol), 9E, 12E, 15E-octadecadien-1-ol (elaidolinolenyl alcohol), 12-hydroxy-9-octadecen-1-ol (ricinoleyl alcohol), 1-eicosanol (arachidyl alcohol) and any combination thereof, wherein the natural oil(s) are selected from the group consisting of straight vegetable oil (SVO), plant parts oils (PPO), tree based oil (TBO), animal fat oil and any combination thereof, and wherein the fuel additive(s) is selected from the group consisting of anti-polymerizing agents, anti-knocking agents, anti-freezing agents, anti-coagulation agents, anti-precipitation agents, cetane index boosters, cetane number boosters and any combination thereof, wherein said fuel additive(s) is present at a concentration ranging from about 0.01 to about 5% v/v.

3. The biofuel composition as claimed in claim 1, wherein the ratio of normal chain alcohol(s) and natural oil(s) is 1:1.

4. The biofuel composition as claimed in claim 1, wherein the biofuel composition comprises normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 5% v/v, natural oil(s) at a concentration of about 5% v/v and diesel at a concentration of about 90% v/v, optionally along with fuel additive(s).

5. The biofuel composition as claimed in claim 1, wherein the biofuel composition comprises normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 7.5% v/v, natural oil(s) at a concentration of about 7.5% v/v and diesel at a concentration of about 85% v/v, optionally along with fuel additive(s).

6. The biofuel composition as claimed in claim 1, wherein the biofuel composition comprises normal chain alcohol(s)

having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 10% v/v, natural oil(s) at a concentration of about 10% v/v and diesel at a concentration of about 80% v/v, optionally along with fuel additive(s).

7. The biofuel composition as claimed in claim 1, wherein the biofuel composition comprises normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 20% v/v, natural oil(s) at a concentration of about 20% v/v and diesel at a concentration of about 60% v/v, optionally along with fuel additive(s).

8. The biofuel composition as claimed in claim 1, wherein the biofuel composition comprises normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 10% v/v, natural oil(s) at a concentration of about 5% v/v and diesel at a concentration of about 85% v/v, optionally along with fuel additive(s).

9. A biofuel composition comprising n-butanol at a concentration ranging from about 1% to about 24% v/v, natural oil(s) at a concentration ranging from about 1% to about 24% v/v and diesel at a concentration ranging from about 52% to about 98% v/v, optionally along with fuel additive(s).

10. A process for preparation of biofuel composition consisting of normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration ranging from about 1% to about 24% v/v, natural oil(s) at a concentration ranging from about 1% to about 24% v/v and diesel at a concentration ranging from about 52% to about 98% v/v, optionally along with fuel additive(s), wherein the natural oil(s) comprise triglycerides, said process comprises mixing said normal chain alcohol(s) with a blend comprising the natural oil(s) and the diesel, optionally along with fuel additive(s).

11. The process as claimed in claim 10, wherein the normal chain alcohols are selected from the group consisting of n-butanol, n-pentanol, n-hexanol, n-heptanol, n-octanol (capryl alcohol), n-nonanol (pelargonic alcohol), n-decanol (capric alcohol), n-dodecanol (lauryl alcohol), n-pentadecanol, n-hexadecanol (cetyl alcohol), n-tetradecanol (myristyl alcohol), cis-9-hexadecen-1-ol (palmitoleyl alcohol), n-octadecanol (stearyl alcohol), 9E-octadecen-1-ol (elaidyl alcohol), cis-9-octadecen-1-ol (oleyl alcohol), 9Z, 12Z-octadecadien-1-ol (linoleyl alcohol), 9E, 12E-octadecadien-1-ol (elaidolinoleyl alcohol), 9Z, 12Z, 15Z-octadecatrien-1-ol (linolenyl alcohol), 9E, 12E, 15E-octadecadien-1-ol (elaidolinolenyl alcohol), 12-hydroxy-9-octadecen-1-ol (ricinoleyl alcohol), 1-eicosanol (arachidyl alcohol) and any combination thereof, wherein the natural oil(s) are selected from a group compris-

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ing straight vegetable oil (SVO), plant parts oils (PPO), tree based oil (TBO), animal fat oil or any combinations thereof, wherein the natural oil(s) are obtained by mechanical or chemical treatment of the source from which it is derived.

12. The process as claimed in claim 10, wherein the ratio of normal chain alcohol(s) and natural oils is 1:1.

13. The process as claimed in claim 10, wherein the fuel additive(s) is selected from the group consisting of anti-polymerizing agents, anti-knocking agents, anti-freezing agents, anti-coagulation agents, anti-precipitation agents, cetane index boosters, cetane number boosters and any combination thereof, wherein the fuel additive(s) is present at a concentration ranging from about 0.01 to about 5% v/v.

14. The process as claimed in claim 10, wherein the process provides for biofuel composition comprising normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 5% v/v, natural oil(s) at a concentration of about 5% v/v and diesel at a concentration of about 90% v/v, optionally along with fuel additive(s).

15. The process as claimed in claim 10, wherein the process provides for biofuel composition comprising normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 7.5% v/v, natural oil(s) at a concentration of about 7.5% v/v and diesel at a concentration of about 85% v/v, optionally along with fuel additive(s).

16. The process as claimed in claim 10, wherein the process provides for biofuel composition comprising normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 10% v/v, natural oil(s) at a concentration of about 10% v/v and diesel at a concentration of about 80% v/v, optionally along with fuel additive(s).

17. The process as claimed in claim 10, wherein the process provides for biofuel composition comprising normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 20% v/v, natural oil(s) at a concentration of about 20% v/v and diesel at a concentration of about 60% v/v, optionally along with fuel additive(s).

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18. The process as claimed in claim 10, wherein the process provides for biofuel composition comprising normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration of about 10%v/v, natural oil(s) at a concentration of about 5% v/v and diesel at a concentration of about 85% v/v, optionally along with fueladditive(s).

19. The process as claimed in claim 10, wherein the process provides for biofuel composition comprising butanol at a concentration ranging from about 1% to about 24% v/v, natural oil(s) at a concentration ranging from about 1% to about 24% v/v and diesel at a concentration ranging from about 52% to about 98% v/v, optionally along with fuel additive(s).

20. A method of fueling biofuel composition consisting of normal chain alcohol(s) having greater than 3 carbon atoms but not more than 21 carbon atoms at a concentration ranging from about 1% to about 24% v/v, natural oil(s) at a concentration ranging from about 1% to about 24% v/v and diesel at a concentration ranging from about 52% to about 98% v/v, optionally along with fuel additive(s), wherein the normal oil(s) comprise triglycerides, said method comprising fueling the biofuel composition for fueling applications.

21. The method of fueling as claimed in claim 20, wherein the fueling applications are selected from the group consisting of space heating, lighting, cooking, running of engines in automobiles, running of motor pumps and running of generators.

22. The biofuel composition of claim 2, wherein the fuel additive(s) are present at a concentration ranging from about 0.01% to about 1% v/v.

23. The process as claimed in claim 13, wherein the fuel additives are present at a concentration ranging from about 0.01% to about 1% v/v.

24. the bifuel composition of claim 1, wherein the ratio of natural oils: normal alcohols is in the range of about 1:1 to about 1:2.

25. the process of claim 10, wherein the ratio of natural oils:normal alcohols is in the range of about 1:1 to about 1:2.

\* \* \* \* \*